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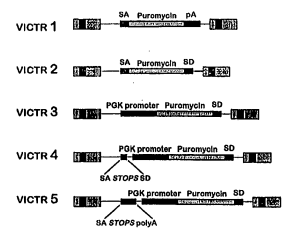
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(54) Title: AN INDEXED LIBRARY OF CELLS CONTAINING GENOMIC MODIFICATIONS AND METHODS OF MAKING AND UTILIZING THE SAME



(57) Abstract

Methods and vectors (both DNA and retroviral) are provided for the construction of a Library of mutated cells. The Library will preferably contain mutations in essentially all genes present in the genome of the cells. The nature of the Library and the vectors allow for methods of screening for mutations in specific genes, and for gathering nucleotide sequence data from each mutated gene to provide a database of tagged gene sequences. Such a database provides a means to access the individual mutant cell clones contained in the Library. The invention includes the described Library, methods of making the same, and vectors used to construct the Library. Methods are also provided for accessing individual parts of the Library either by sequence or by pooling and screening. The invention also provides for the generation of non-human transgenic animals which are mutant for specific genes as isolated and generated from the cells of the Library.

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AN INDEXED LIBRARY OF CELLS CONTAINING GENOMIC MODIFICATIONS AND METHODS OF MAKING AND UTILIZING THE SAME

The present application claims priority to U.S.

5 Applications Ser. Nos. 08/726,867, filed October 4, 1996, 08/728,963, filed October 11, 1996, and 08/907,598, filed August 8, 1997, the disclosures of which are herein incorporated by reference.

1.0. FIELD OF THE INVENTION

The invention relates to an indexed library of genetically altered cells and methods of organizing the cells into an easily manipulated and characterized Library. The invention also relates to methods of making the library, vectors for making insertion mutations in genes, methods of gathering sequence information from each member clone of the Library, and methods of isolating a particular clone of interest from the Library.

20 2.0. BACKGROUND OF THE INVENTION

The general technologies of targeting mutations into the genome of cells, and the process of generating mouse lines from genetically altered embryonic stem (ES) cells with specific genetic lesions are well known (Bradley, 1991, Cur. Opin. Biotech. 2:823-829). A random method of generating genetic lesions in cells (called gene, or promoter, trapping) has been developed in parallel with the targeted methods of genetic mutation (Allen et al., 1988 Nature 333(6176):852-855; Brenner et al., 1989, Proc. Natl. Acad. Sci. U.S.A. 30 86(14):5517-5521; Chang et al., 1993, Virology 193(2):737-747; Friedrich and Soriano, 1993, Insertional mutagenesis by retroviruses and promoter traps in embryonic stem cells, p. 681-701. In Methods Enzymol., vol. 225., P. M. Wassarman and M. L. DePamphilis (ed.), Academic Press, Inc., San Diego; 35 Friedrich and Soriano, 1991, Genes Dev. 5(9):1513-1523; Gossler et al., 1989, Science 244(4903):463-465; Kerr et al., 1989, Cold Spring Harb. Symp. Quant. Biol. 2:767-776; Reddy

et al., 1991, J Virol. 65(3):1507-1515; Reddy et al., 1992,

Proc. Natl. Acad. Sci. U.S.A. 89(15):6721-6725; Skarnes et al., 1992, Genes Dev. 6(6):903-918; von Melchner and Ruley, 1989, J. Virol. 63(8):3227-3233; Yoshida et al., 1995, Transgen. Res. 4:277-287). Gene trapping provides a means to 5 create a collection of random mutations by inserting fragments of DNA into transcribed genes. Insertions into transcribed genes are selected over the background of total insertions since the mutagenic DNA encodes an antibiotic resistance gene or some other selectable marker. 10 selectable marker lacks its own promoter and enhancer and must be expressed by the endogenous sequences that flank the marker after it has integrated. Using this approach, transcription of the selectable marker is activated and the cell gene is concurrently mutated. This type of strict 15 selection makes it possible to easily isolate thousands of ES cell colonies, each with a unique mutagenic insertion.

Collecting mutants on a large-scale has been a powerful genetic technique commonly used for organisms which are more amenable to such analysis than mammals. These organisms,

- 20 such as Drosophila melanogastor, yeast Saccharomyces cerevisiae, and plants such as Arabadopsis thalia are small, have short generation times and small genomes (Bellen et al., 1989, Genes Dev. 3(9):1288-1300; Bier et al., 1989, Genes Dev. 3(9):1273-1287; Hope, 1991, Develop. 113(2):399-408.
- 25 These features allow an investigator to rear many thousands or millions of different mutant strains without requiring unmanageable resources. However, these type of organisms have only limited value in the study of biology relevant to human physiology and health. It is therefore important to
- 30 have the power of large-scale genetic analysis available for the study of a mammalian species that can aid in the study of human disease. Given that the entire human genome is presently being sequenced, the comprehensive genetic analysis of a related mammalian species will provide a means to
- 35 determine the function of genes cloned from the human genome. At present, rodents, and particularly mice, provide the best model for genetic manipulation and analysis of mammalian

physiology.

Gene trapping has been used as an analytical tool to identify genes and regulatory regions in a variety of animal cell types. One system that has proved particularly useful is based on the use of ROSA (reverse orientation splice acceptor) retroviral vectors (Friedrich and Soriano, 1991 and 1993).

The ROSA system can generate mutations that result in a detectable homozygous phenotype with a high frequency. About 10 50% of all the insertions caused embryonic lethality. The specifically mutated genes may easily be cloned since the gene trapping event produces a fusion transcript. This fusion transcript has trapped exon sequences appended to the sequences of the selectable marker allowing the latter to be 15 used as a tag in polymerase chain reaction (PCR)-based protocols, or by simple cDNA cloning. Examples of genes isolated by these methods include a transcription factor related to human TEF-1 (transcription enhancer factor-1) which is required in the development of the heart (Chen et 20 al., 1994, Genes Devel. 8:2293-2301. Another (spock), is distantly related to yeast genes encoding secretion proteins and is important during gastrulation.

The above experiments have established that the ROSA system is an effective analytical tool for genetic analysis 25 in mammals. However, the structure of many ROSA vectors selects for the "trapping" of 5' exons which, in many cases, do not encode proteins. Such a result is adequate where one wishes to identify and eventually clone control (i.e., promoter or enhancer) sequences, but is not optimal where the 30 generation of insertion-inactivated null mutations is desired, and relevant coding sequence is needed. Thus, the construction of large-scale mutant (preferably null mutant) libraries requires the use of vectors that have been designed to select for insertion events that have occurred within the 35 coding region of the mutated genes as well as vectors that are not limited to detecting insertions into expressed genes.

3.0. SUMMARY OF THE INVENTION

An object of the present invention is to provide a set of genetically altered cells (the 'Library'). The genetic alterations are of sufficient randomness and frequency such 5 that the combined population of cells in the Library represent mutations in essentially every gene found in the cell's genome. The Library is used as a source for obtaining specifically mutated cells, cell lines derived from the individually mutated cells, and cells for use in the 10 production of transgenic non-human animals.

A further object is to provide the vectors, both DNA and retroviral based, that may be used to generate the Library. Typically, at least two distinct vector designs will be used in order to mutate genes that are actively expressed in the target cell, and genes that are not expressed in the target cell. Combining the mutant cells obtained using both types of vectors best ensures that the Library provides a comprehensive set of gene mutations.

A particularly useful vector class contemplated by the 20 present invention includes a vector for inserting foreign exons into animal cell transcripts that comprises a selectable marker, a promoter element operatively positioned 5' to the selectable marker, a splice donor site operatively positioned 3' to the selectable marker, and a second 25 mutagenic foreign polynucleotide sequence located upstream from the promoter element that disrupts, or otherwise "poisons", the splicing or read-through expression of the endogenous cellular transcript. Typically, the mutagenic foreign polynucleotide sequence may incorporate a 30 polyadenylation (pA) site, a nested set of stop codons in each of the three reading frames, splice acceptor and splice donor sequences in operable combination, a mutagenic exon, or any mixture of mutagenic features that effectively prevent the expression of the cellular gene. For example, a 35 polyadenylation sequence may be incorporated in addition to or in lieu of the splice donor sequence. A preferred organization for the mutagenic polynucleotide sequence

comprises a polyadenylation site positioned upstream from a selectable marker which is in turn located upstream from a splice acceptor sequence. Preferably, such a vector does not comprise a transcription terminator or polyadenylation site operatively positioned relative to the coding region of the selectable marker, and shall not comprise a splice acceptor site operatively positioned between the promoter element and the initiation codon of said selectable marker.

An additional vector contemplated by the present

10 invention is designed to replace the normal 3' end of an animal cell transcript with a foreign exon. Such a vector shall generally be engineered to comprise a selectable marker, a splice acceptor site operatively positioned upstream (5') from the initiation codon of the selectable

15 marker, and a polyadenylation site operatively positioned downstream (3') from the termination codon (3' end) of the selectable marker. Preferably, the vector will not comprise a promoter element operatively positioned upstream from the coding region of the selectable marker, and will not comprise a splice donor sequence operatively positioned between the 3' end of the coding region of the selectable marker and the polyadenylation site.

Yet another vector contemplated by the present invention is a vector designed to insert a mutagenic foreign

25 polynucleotide sequence within an animal cell transcript (i.e., the foreign polynucleotide sequence is flanked on both sides by endogenous exons). As described above, the mutagenic foreign polynucleotide sequence may be any sequence that disrupts the normal expression of the gene into which

30 the vector has integrated. Optionally, the vector may additionally incorporate a selectable marker, a splice acceptor site operatively positioned 5' to the initiation codon of the selectable marker, a splice donor site operatively positioned 3' to said selectable marker.

35 Preferably, this vector shall not comprise a polyadenylation site operatively positioned 3' to the coding region of said selectable marker, and shall not comprise a promoter element

operatively positioned 5' to the coding region of said selectable marker.

An additional embodiment of the present invention is a library of genetically altered cells that have been treated 5 to stably incorporate one or more types of the vectors The presently described library of described above. cultured animal cells may be made by a process comprising the steps of treating (i.e., infecting, transfecting, retrotransposing, or virtually any other method of 10 introducing polynucleotides into a cell) a population of cells to stably integrate a vector that mediates the splicing of a foreign exon internal to a cellular transcript, transfecting another population of cells to stably integrate a vector that mediates the splicing of a foreign exon 5' to 15 an exon of a cellular transcript, and selecting for transduced cells that express the products encoded by the foreign exons.

Alternatively, an additional embodiment of the present invention describes a mammalian cell library made by a method comprising the steps of: transfecting a population of cells with a vector capable of expressing a selectable marker in the cell only after the vector inserts into the host genome; transfecting or infecting a population of cells with a vector containing a selectable marker that is substantially only expressed by cellular control sequences (after the vector integrates into the host cells genome); and growing the transfected cells under conditions that select for the expression of the selectable marker.

In an additional embodiment of the present invention,
30 the two populations of transfected cells will be individually
grown under selective conditions, and the resulting mutated
population of cells collectively comprises a substantially
comprehensive library of mutated cells.

In an additional embodiment of the present invention,

35 the individual mutant cells in the library are separated and clonally expanded. Additionally, the clonally expanded mutant cells may then be analyzed to ascertain the DNA

sequence, or partial DNA sequence of the mutated host gene.

The presently described methods of making, organizing, and indexing libraries of mutated animal cells are also broadly applicable to virtually any eukaryotic cells that may be genetically manipulated and grown in culture.

The invention provides for sequencing every gene mutated in the Library. The resulting sequence database subsequently serves as an index for the library. In essence, every cell line in the Library is individually catalogued using the 10 partial sequence information. The resulting sequence is specific for the mutated gene since the present methods are designed to obtain sequence information from exons that have been spliced to the marker sequence. Since the coverage of the mutagenesis is preferably the entire set of genes in the 15 genome, the resulting Library sequence database contains sequence from essentially every gene in the cell. From this database, a gene of interest can be identified. Once identified, the corresponding mutant cell may be withdrawn from the Library based on cross reference to the sequence 20 data.

An additional embodiment of the invention provides for methods of isolating mutations of interest from the Library. Two methods are proposed for obtaining individual mutant cell lines from the Library. The first provides a scheme where clones of the cells generated using the above vectors are pooled into sets of defined size. Using the procedure described below which utilizes reverse transcription (RT) and polymerase chain reaction (PCR), a cell line with a mutation in a gene whose sequence is partly or wholly known is isolated from organized sets of these pools. A few rounds of this screening procedure results in the isolation of the desired individual cell line.

A second procedure involves the sequencing of regions flanking the vector insertion sites in the various cells in the library. The sequence database generated from these data effectively constitutes an index of the clones in the library that may be used to identify cells having mutations in

specific genes.

4.0. DESCRIPTION OF THE FIGURES

- Figure 1. Shows a diagrammatic representation of 5 different 5 vectors that are generally representative of the type of vectors that may be used in the present invention.
- Figure 2. Shows a general strategy for identifying "trapped" cellular sequences by PCR analysis of the cellular exons that 10 flank the foreign intron introduced by the VICTR 2 vector.
 - Figure 3 shows a PCR based strategy for identifying tagged genes by chromosomal location.
- 15 Figure 4. Is a diagrammatic representation of a strategy of identifying or indexing the specific clones in the library via PCR analysis and sequencing of mRNA samples obtained from the cells in the library.
- 20 Figure 5. Is a diagrammatic representation of a method of isolating positive clones by screening pooled mutant cell clones.
- Figure 6. Partial nucleic acid or predicted amino acid 25 sequence data from 9 clones (OST1-9) isolated using the described techniques aligned with similar sequences from previously characterized genes.
- Figure 7. Provides a diagrammatic representation of VICTRs 3 30 and 20 as well as the transcripts that result after integration into a hypothetical region of the target cell genome (i.e., "Wildtype Locus).
- Figure 8. Provides a representative list of a portion of the 35 known genes that have been identified using the disclosed methods and technology.

5.0. DETAILED DESCRIPTION OF THE INVENTION

The present invention describes a novel indexed library containing a substantially comprehensive set of mutations in the host cell genome, and methods of making and using the 5 same. The presently described Library comprises as a set of cell clones that each possess at least one mutation (and preferably a single mutation) caused by the insertion of DNA that is foreign to the cell. For the purposes of the present invention, "foreign" polynucleotide sequences can be any 10 sequences that are newly introduced to a cell, do not naturally occur in the cell at the engineered region of the chromosome, or occur in the cell but are not organized to provide an identical function to that provided in the engineered vector.

The particularly novel features of the Library include 15 the methods of construction, and indexing. To index the library, the mutant cells of the library are clonally expanded and each mutated gene is at least partially sequenced. The Library thus provides a novel tool for 20 assessing the specific function of a given gene. The insertions cause a mutation which allow for essentially every gene represented in the Library to be studied using genetic techniques either in vitro or in vivo (via the generation of For the purposes of the present transgenic animals). 25 invention, the term "essentially every gene" shall refer to the statistical situation where there is generally at least about a 70 percent probability that the genomes of cells used to construct the library collectively contain at least one inserted vector sequence in each gene, preferably a 85 30 percent probability, and more specifically at least about a 95 percent probability as determined by a standard Poisson distribution.

Also for the purposes of the present invention the term "gene" shall refer to any and all discrete coding regions of 35 the cell's genome, as well as associated noncoding and regulatory regions. Additionally, the term operatively positioned shall refer to the control elements or genes that

are provided with the proper orientation and spacing to provide the desired or indicated functions of the control elements or genes.

For the purposes of the present invention, a gene is

5 "expressed" when a control element in the cell mediates the
production of functional or detectable levels of mRNA encoded
by the gene, or a selectable marker inserted therein. A gene
is not expressed where the control element in the cell is
absent, has been inactivated, or does not mediate the
production of functional or detectable levels of mRNA encoded
by the gene, or a selectable marker inserted therein.

5.1. Vectors used to build the Library

A number of investigators have developed gene trapping
15 vectors and procedures for use in mouse and other cells
(Allen et al., 1988; Bellen et al., 1989, Genes Dev.
3(9):1288-1300; Bier et al., 1989, Genes Dev. 3(9):1273-1287;
Bonnerot et al., 1992, J Virol. 66(8):4982-4991; Brenner et
al., 1989; Chang et al., 1993; Friedrich and Soriano, 1993;

- 20 Friedrich and Soriano, 1991; Goff, 1987, Methods Enzymol.
 152:469-481; Gossler et al.; Hope, 1991; Kerr et al., 1989;
 Reddy et al., 1991; Reddy et al., 1992; Skarnes et al., 1992;
 von Melchner and Ruley; Yoshida et al., 1995). The gene
 trapping system described in the present invention is based
- 25 on significant improvements to the published SA (splice acceptor) DNA vectors and the ROSA (reverse orientation, splice acceptor) retroviral vectors (Chen et al., 1994; Friedrich and Soriano, 1991 and 1993). The presently described vectors also use a selectable marker called βgeo.
- 30 This gene encodes a protein which is a fusion between the β -galactosidase and neomycin phosphotransferase proteins. The presently described vectors place a splice acceptor sequence upstream from the βgeo gene and a poly-adenylation signal sequence downstream from the marker. The marker is
- 35 integrated after transfection by, for example, electroporation (DNA vectors), or retroviral infection, and gene trap events are selected based on resistance to G418

resulting from activation of βgeo expression by splicing from the endogenous gene into the ROSA splice acceptor. This type of integration disrupts the transcription unit and preferably results in a null mutation at the locus.

tool, the present invention contemplates gene trapping on a large scale. The vectors utilized in the present invention have been engineered to overcome the shortcomings of the early gene trap vector designs, and to facilitate procedures allowing high throughput. In addition, procedures are described that allow the rapid and facile acquisition of sequence information from each trapped cDNA which may be adapted to allow complete automation. These latter procedures are also designed for flexibility so that additional molecular information can easily be obtained subsequently. The present invention therefore incorporates gene trapping into a larger and unique tool. A specially organized set of gene trap clones that provide a novel and powerful new tool of genetic analysis.

The presently described vectors are superficially 20 similar to the ROSA family of vectors, but constitute significant improvements and provide for additional features that are useful in the construction and indexing of the Library. Typically, gene trapping vectors are designed to 25 detect insertions into transcribed gene regions within the genome. They generally consist of a selectable marker whose normal expression is handicapped by exclusion of some element required for proper transcription. When the vector integrates into the genome, and acquires the necessary 30 element by juxtaposition, expression of the selectable marker is activated. When such activation occurs, the cell can survive when grown in the appropriate selective medium which allows for the subsequent isolation and characterization of the trapped gene. Integration of the gene trap generally 35 causes the gene at the site of integration to be mutated.

Some gene trapping vectors have a splice acceptor preceding a selectable marker and a poly-adenylation signal

following the selectable marker, and the selectable marker gene has its own initiator ATG codon. Using this arrangement, the fusion transcripts produced after integration generally only comprise exons 5' to the insertion 5 site to the known marker sequences. Where the vector has inserted into the 5' region of the gene, it is often the case that the only exon 5' to the vector is a non-coding exon. Accordingly, the sequences obtained from such fusions do not provide the desired sequence information about the relevant 10 gene products. This is because untranslated sequences are generally less well conserved than coding sequences.

To compensate for the short-comings of earlier vectors, the vectors of the present invention have been designed so that 3' exons are appended to the fusion transcript by 15 replacing the poly-adenylation and transcription termination signals of earlier ROSA vectors with a splice donor (SD) sequence. Consequently transcription and splicing generally results in a fusion between all or most of the endogenous transcript and the selectable marker exon, for example βgeo , 20 neomycin (neo) or puromycin (puro). The exon sequences immediately 3' to the selectable marker exon may then be sequenced and used to establish a database of expressed sequence tags. The presently described procedures will typically provide approximately 200 nucleotides of sequence, These sequences will generally be coding and therefore informative. The prediction that the sequence obtained will be from coding region is based on two factors. First, gene trap vectors are generally found near the 5' end of the gene immediately after untranslated exons because the 30 method selects for integration events that place the initiator ATG of the selectable marker as the first encountered, and thus used, for translation. Second, mammalian transcripts have short 5' untranslated regions (UTRs) which are typically between 50 and 150 nucleotides in 35 length.

The obtained sequence information also provides a ready source of probes that may be used to isolate the full-length

gene or cDNA from the host cell, or as heterologous probes for the isolation of homologous genes in other species.

Internal exons in mammalian transcripts are generally quite small, on the average 137 bases with few over 300 5 bases. Consequently, a large internal exon may be spliced less efficiently. Thus, the presently described vectors have been designed to sandwich relatively small selectable markers (for example: neo ,~800 bases, or a smaller drug resistance gene such as puro ,~600 bases) between the requisite splicing 10 elements to produce relatively small exons. Exons of this size are more typical of mammalian exons and do not present undue problems for the splicing machinery of the cell. a design consideration is novel to the presently disclosed gene trapping vectors. Accordingly, an additional embodiment 15 of the claimed vectors is that the respective splice acceptor and splice donor sites are engineered such that they are operatively positioned close to the ends of the selectable marker coding region (the region spanning from the initiation codon to the termination codon). Generally, the splice 20 acceptor or splice donor sequences shall appear within about 80 bases from the nearest end of the selectable marker coding region, preferably within about 50 bases from the nearest end of the coding region, more preferably within about 30 bases from the nearest end of the coding regions and specifically 25 within about 20 bases of the nearest end of the selectable marker coding region.

The new vectors are represented in retroviral form in Figure 1. They are used by infecting target cells with retroviral particles such that the proviruses shown in the 30 schematic can be found in the genome of the target. These vectors are called VICTR which is an acronym for "viral constructs for trapping".

The presently described retroviral vectors may be used in conjunction with retroviral packaging cell lines such as 35 those described in U.S. Patent No. 5,449,614 ("'614 patent") issued September 12, 1995, herein incorporated by reference. Where non-mouse animal cells are to be used as targets for

generating the described libraries, packaging cells producing retrovirus with amphotropic envelopes will generally be employed to allow infection of the host cells.

The mutagenic gene trap DNA may also be introduced into the target cell genome by various transfection techniques which are familiar to those skilled in the art such as electroporation, lipofection, calcium phosphate precipitation, infection, retrotransposition, and the like. Examples of such techniques may be found in Sambrook et al. (1989) Molecular Cloning Vols. I-III, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York, and Current Protocols in Molecular Biology (1989) John Wiley & Sons, all Vols. and periodic updates thereof, herein incorporated by reference. The transfected versions of the retroviral vectors are typically plasmid DNA molecules containing DNA cassettes comprising the described features between the retroviral LTRs.

The vectors VICTR 1 and 2 (Fig. 1) are designed to trap genes that are transcribed in the target cell. To trap genes 20 that are not expressed in the target cell, gene trap vectors such as VICTR 3, 4 and 5 (described below) are provided. These vectors have been engineered to contain a promoter element capable of initiating transcription in virtually any cell type which is used to transcribe the coding sequence of 25 the selectable marker. However, in order to get proper translation of the marker product, and thus render the cell resistant to the selective antibiotic, a polyadenylation signal and a transcription termination sequence must be provided. Vectors VICTR 3 through 5 are constructed such 30 that an effective polyadenylation signal can only be provided by splicing with an externally provided downstream exon that contains a poly-adenylation site. Therefore, since the selectable marker coding region ends only in a splice donor sequence, these vectors must be integrated into a gene in 35 order to be properly expressed. In essence, these vectors append the foreign exon encoding the marker to the 5' end of an endogenous transcript. These events will tag genes and

create mutations that are used to make clones that will become part of the Library.

With the above design considerations, the VICTR series of vectors, or similarly designed and constructed vectors, 5 have the following features. VICTR 1 is a terminal exon gene trap. VICTR 1 does not contain a control region that effectively mediates the expression of the selectable marker gene. Instead, the coding region of the selectable marker contained in VICTR 1, in this case encoding puromycin 10 resistance (but which can be any selectable marker functional in the target cell type), is preceded by a splice acceptor sequence and followed by a polyadenylation addition signal sequence. The coding region of the puro gene has an initiator ATG which is downstream and adjacent to a region of 15 sequence that is most favorable for translation initiation in eukaryotic cells - the so called Kozak consensus sequence (Kozak, 1989, J. Cell, Biol. 108(2):229-241). With a Kozak sequence and an initiator ATG, the puro gene in VICTR 1 is activated by integrating into the intron of an active gene, 20 and the resulting fusion transcript is translated beginning at the puromycin initiation (ATG/AUG) codon. However, terminal gene trap vectors need not incorporate an initiator ATG codon. In such cases, the gene trap event requires splicing and the translation of a fusion protein that is 25 functional for the selectable marker activity. The inserted puromycin coding sequence must therefore be translated in the same frame as the "trapped" gene.

The splice acceptor sequence used in VICTR 1 and other members of the VICTR series is derived from the adenovirus 30 major late transcript splice site located at the intron 1/exon 2 boundary. This sequence contains a polypyrimidine stretch preceding the AG dinucleotide which denotes the actual splice site. The presently described vectors contemplate the use of any similarly derived splice acceptor sequence. Preferably, the splice acceptor site will only rarely, if ever, be involved in alternative splicing events.

The polyadenylation signal at the end of the puro gene is derived from the bovine growth hormone gene. Any similarly derived polyadenylation signal sequence could be used if it contains the canonical AATAAA and can be demonstrated to terminate transcription and cause a polyadenylate tail to be added to the engineered coding exons.

VICTR 2 is a modification of VICTR 1 in which the polyadenylation signal sequence is removed and replaced by a 10 splice donor sequence. Like VICTR 1, VICTR 2 does not contain a control region that effectively mediates the expression of the selectable marker gene. Typically, the splice donor sequence to be employed in a VICTR series vector shall be determined by reference to established literature or 15 by experimentation to identify which sequences properly initiate splicing at the 5' end of introns in the desired target cell. The specifically exemplified sequence, AGGTAAGT, results in splicing occurring in between the two G bases. Genes trapped by VICTR 2 splice upstream exons onto 20 the puro exon and downstream exons onto the end of the puro exon. Accordingly, VICTR 2 effectively mutates gene expression by inserting a foreign exon in-between two naturally occurring exons in a given transcript. Again, the puro gene may or may not contain a consensus Kozak 25 translation initiation sequence and properly positioned ATG initiation codon. As discussed above, gene trapping by VICTR 1 and VICTR 2 requires that the mutated gene is expressed in the target cell line. By incorporating a splice donor into the VICTR traps, transcript sequences downstream 30 from the gene trap insertion can be determined. As described above, these sequences are generally more informative about the gene mutated since they are more likely to be coding sequences. This sequence information is gathered according to the procedures described below.

VICTR 3, VICTR 4 and VICTR 5 are gene trap vectors that do not require the cellular expression of the endogenous trapped gene. The VICTR vectors 3 through 5 all comprise a

promoter element that ensures that transcription of the selectable marker would be found in all cells that have taken up the gene trap DNA. This transcription initiates from a promoter, in this case the promoter element from the mouse 5 phosphoglycerate kinase (PGK) gene. However, since the constructs lack a polyadenylation signal there can be no proper processing of the transcript and therefore no translation. The only means to translate the selectable marker and get a resistant cell clone is by acquiring a 10 polyadenylation signal. Since polyadenylation is known to be concomitant with splicing, a splice donor is provided at the end of the selectable marker. Therefore, the only positive gene trap events using VICTR 3 through 5 will be those that integrate into a gene's intron such that the marker exon is 15 spliced to downstream exons that are properly polyadenylated. Thus genes mutated with the VICTR vectors 3 through 5 need not be expressed in the target cell, and these gene trap vectors can mutate all genes having at least one intron. The design of VICTR vectors 3 through 5 requires a promoter 20 element that will be active in the target cell type, a Although a selectable marker and a splice donor sequence. specific promoter was used in the specific embodiments, it should be understood that appropriate promoters may be selected that are known to be active in a given cell type. 25 Typically, the considerations for selecting the splice donor sequence are identical to those discussed for VICTR 2, supra. VICTR 4 differs from VICTR 3 only by the addition of a small exon upstream from the promoter element of VICTR 4. This exon is intended to stop normal splicing of the mutated 30 gene. It is possible that insertion of VICTR 3 into an intron might not be mutagenic if the gene can still splice between exons, bypassing the gene trap insertion. The exon

in VICTR 4 is constructed from the adenovirus splice acceptor described above and the synthetic splice donor also described 35 above. Stop codons are placed in all three reading frames in the exon, which is about 100 bases long. The stops would truncate the endogenous protein and presumably cause a

mutation.

A conceptually similar alternative design uses a terminal exon like that engineered into VICTR 5. Instead of a splice donor, a polyadenylation site is used to terminate 5 transcription and produce a truncated message. Stops in all three frames are also provided to truncate the endogenous protein as well as the resulting transcript.

VICTR 20 is a modified version of VICTR 3 that incorporates a polyadenylation site 5' to the PGK promoter, 10 the IRESβgeo sequence (i.e., foreign mutagenic polynucleotide sequence) 5' to the polyadenylation site, and a splice acceptor site 5' to the IRESβgeo coding region. VICTR 20 additionally incorporates, in operable combination, a pair of recombinase recognition sites that flank the PGKpuroSD 15 cassette.

All of the traps of the VICTR series are designed such that a fusion transcript is formed with the trapped gene. For all but VICTR 1, the fusion contains cellular exons that are located 3' to the gene trap insertion. All of the 20 flanking exons may be sequenced according to the methods described in the following section. To facilitate sequencing, specific sequences are engineered onto the ends of the selectable marker (e.g., puromycin coding region). Examples of such sequences include, but are not limited to 25 unique sequences for priming PCR, and sequences complementary to the standard M13 forward sequencing primer. Additionally, stop codons are added in all three reading frames to ensure that no anomalous fusion proteins are produced. All of the unique 3' primer sequences are followed immediately by the 30 synthetic 9 base pair splice donor sequence. This keeps the size of the exon comprising the selectable marker (puro gene) at a minimum to best ensure proper splicing, and positions the amplification and sequencing primers immediately adjacent to the flanking "trapped" exons to be sequenced as part of 35 the construction of a Library database.

When any members of the VICTR series are constructed as retroviruses, the direction of transcription of the

selectable marker is opposite to that of the direction of the normal transcription of the retrovirus. The reason for this organization is that the transcription elements such as the polyadenylation signal, the splice sites and the promoter elements found in the various members of the VICTR series interfere with the proper transcription of the retroviral genome in the packaging cell line. This would eliminate or significantly reduce retroviral titers. The LTRs used in the construction of the packaging cell line are self-

10 inactivating. That is, the enhancer element is removed from the 3' U3 sequences such that the proviruses resulting from infection would not have an enhancer in either LTR. An enhancer in the provirus may otherwise affect transcription of the mutated gene or nearby genes.

Since a 'cryptic' splice donor sequence is found in the inverted LTRs, this splice donor sequence has been removed from the VICTR vectors by site specific mutagenesis. It was deemed necessary to remove this splice donor so that it would not affect the trapping splicing events.

The present disclosure also describes vectors that 20 incorporate a new way to conduct positive selection. VICTR 3 and VICTR 20 are two examples of such vectors. Both VICTR 3 and VICTR 20, contain PGKpuroSD which must splice into exons of gene that provide a polyadenylation addition sequence in 25 order to allow expression of the puromycin selectable marker gene. When placed in a targeting vector, PGKpuroSD allows for positive selection when targeting takes place. addition to providing positive selection, targeted events among resistant colonies are easy to identify by the 3' RACE 30 protocols (see section 5.2.2., infra) used for Omnibank production. This automated process allows for the rapid identification of targeted events. It is important that unlike $SA\beta$ geo, PGKpuroSD does not require expression of the targeted gene in order to provide positive selection. 35 addition, VICTR 20 provides 2 potential positive selectable markers (puro and neo). The use of two selectable markers,

when a gene is expressed, provides a means to increase the

targeting efficiency by requiring both selectable markers to function which is much more remote a possibility than having one selectable marker function unless there is a targeted event. The addition of a negative selection cassette to these vectors would only increase their targeting efficiency.

An additional feature that may be incorporated into the presently described vectors includes the use of recombinase recognition sequences. Bacteriophage P1 Cre recombinase and flp recombinase from yeast plasmids are two examples of 10 site-specific DNA recombinase enzymes which cleave DNA at specific target sites (loxP sites for cre recombinase and frt sites for flp recombinase) and catalyze a ligation of this DNA to a second cleaved site. When a piece of DNA is flanked by 2 loxP or frt sites (e.g., recombinase control elements) 15 in the same orientation, the corresponding recombinase will cause the removal of the intervening DNA sequence. piece of DNA is flanked by loxP or frt sites in an indirect orientation, the corresponding recombinase will essentially activate the control elements to cause the intervening DNA to 20 be flipped into the opposite orientation. These recombinases provide powerful approaches for manipulating DNA in situ.

Recombinases have important applications for gene trapping and the production of a library of trapped genes. When constructs containing PGKpuroSD are used to trap genes, 25 the fusion transcript between puromycin and sequences of the trapped gene could result in some level of protein expression from the trapped gene if translational reinitiation occurs. Another important issue is that several reports suggest that the PGK promoter can affect the expression of nearby genes. 30 These effects may make it difficult to determine gene function after a gene trap event since one could not discern whether a given phenotype is associated with the inactivation of a gene, or the transcription of nearby genes. Both potential problems are solved by exploiting recombinase 35 activity. When PGKpuroSD is flanked by loxP, frt, or any other recombinase sites in the same orientation, the addition of the corresponding recombinase will result in the removal

of PGKpuroSD. In this way, effects caused by PGKpuroSD fusion transcripts, or the PGK promoter, are avoided.

Accordingly, a vector that may be particularly useful for the practice of the present invention is VICTR 20. 5 vector replaces the terminal exon of VICTR 5 with a splice acceptor located upstream from the β geo gene which can be used for both LacZ staining and antibiotic selection. fusion gene possesses its own initiator methionine and an internal ribosomal entry site (IRES) for efficient 10 translation initiation. In addition, the PGK promoter and puromycin-splice donor sequences have been flanked by lox P recombination sites. This allows for the ability to both remove and introduce sequences at the integration site and is of potential value with regard to the manipulation of regions 15 proximal to trapped target genes (Barinaga, Science 265:26-8, 1994). While this particular vector includes lox P recombination sites, the present invention is in no way limited to the use of this specific recombination site (Akagi et al., Nucleic Acids Res 25:1766-73, 1997).

Another very important use of recombinases is to produce 20 mutations that can be made tissue-specific and/or inducible. In the presently described vectors, the Sa β geo or SAIRES β geo component provides the mutagenic function by "trapping" the normal splicing from preceding exons. If the $SA\beta$ geo is 25 flanked by inverted loxP, frt, or any other recombinase sites, the addition of the corresponding recombinase results in the flipping of the $SA\beta$ geo sequence so that it no longer prevents the normal splicing of the cellular gene into which it is integrated. To make a gene trap tissue-specific or 30 inducible one could produce the trap with $SA\beta$ geo in the reverse orientation and then provide recombinase activity only at the time and place where one wishes to remove the gene function. The use of tissue-specific or inducible recombinase constructs allows one to choose when and where 35 one removes, or activates, the function of the targeted gene.

One method for practicing the inducible forms of recombinase mediated gene expression involves the use of

vectors that use inducible or tissue specific promoter/operator elements to express the desired recombinase activity. The inducible expression elements are preferably operatively positioned to allow the inducible control or 5 activation of expression of the desired recombinase activity. Examples of such inducible promoters or control elements include, but are not limited to, tetracycline, metallothionine, ecdysone, and other steroid-responsive promoters, rapamycin responsive promoters, and the like (No 10 et al., Proc Natl Acad Sci USA <u>93</u>:3345-51, 1996; Furth et al., Proc Natl Acad Sci USA 91:9302-6, 1994). Additional control elements that can be used include promoters requiring specific transcription factors such as viral, particularly HIV, promoters. Vectors incorporating such promoters would 15 only express recombinase activity in cells that express the necessary transcription factors.

The incorporation of recombinase sites into the gene trapping vectors highlights the value of using the described gene trap vectors to deliver specific DNA sequence elements throughout the genome. Although a variety of vectors are

available for placing sequences into the genome, the presently described vectors facilitate both the insertion of the specific elements, and the subsequent identification of where sequence has inserted into the cellular chromosome.

25 Additionally, the presently described vectors may be used to place recombinase recognition sites throughout the genome. The recombinase recognition sites could then be used to either remove or insert specific DNA sequences at predetermined locations.

Moreover, the described gene trap vectors can also be used to insert regulatory elements throughout the genome.

Recent work has identified a number of inducible or repressible systems that function in the mouse. These include the rapamycin, tetracycline, ecdysone,

35 glucocorticoid, and heavy metal inducible systems. These systems typically rely on placing DNA elements in or near a promoter. An inducible or repressible transcription factor

that can identify and bind to the DNA element may also be engineered into the cells. The transcription factor will specifically bind to the DNA element in either the presence or absence of a ligand that binds to the transcription factor and, depending on the structure of the transcription factor, it will either induce or repress the expression of the cellular gene into which the DNA elements have been inserted. The ability to place these inducible or repressible elements throughout the genome would increase the value of the library 10 by adding the potential to regulate the expression of the trapped gene.

The vectors described also have important applications for the overexpression of genes or portions of genes to select for phenotypic effects. Currently, overexpression of 15 cDNA libraries to look for genes or parts of genes with specific functions is a common practice. One example would be to overexpress genes or portions of genes to look for expression that causes loss of contact inhibition for cell growth as determined by growth in soft agar. This would 20 allow the identification of genes or portions of genes that can act as oncogenes. Simple modifications of VICTR 20 would allow it to be used for these applications. For example, the addition of an internal ribosome entry site (IRES) 3' to the puromycin selectable marker and before the SD sequence, would 25 result in the overexpression of sequences from the trapped downstream exons. In addition, the IRES could be modified by, for example, the addition of one or two nucleotides such that there could be 3 basic vectors that would allow expression of trapped exons in all three reading frames. 30 this way, genes could be trapped throughout the genome resulting in overexpression of genes, or portions thereof, to examine the cellular function of the trapped genes. identification of function could be done by selecting for the function of interest (i.e., growth in soft agar could result 35 from the overexpression of potentially oncogenic genes). This technique would allow for the screening or selection of large numbers of genes, or portions thereof, by

overexpressing the genes and identifying cells displaying the phenotypes of interest. Additional assays could, for example, identify candidate tumor suppressor genes based on their ability, when overexpressed, to prevent growth in soft agar.

Given the fact that expression pattern information can provide insight into the possible functions of genes mutated by the current methods, another LTR vector, VICTR 6, has been constructed in a manner similar to VICTR 5 except that the 10 terminal exon has been replaced with either a gene coding for β -galactosidase (β gal) or a fusion between β -gal and neomycin phosphotransferase (β geo), each proceeded by a splice acceptor and followed by a polyadenylation signal. Endogenous gene expression and splicing of these markers into 15 cellular transcripts and translation into fusion proteins will allow for increased mutagenicity as well as the delineation of expression through Lac Z staining.

An additional vector, VICTR 12, incorporates two separate selectable markers for the analysis of both 20 integration sites and trapped genes. One selectable marker (e.g. puro) is similar to that for VICTRs 3 through 5 in that it contains a promoter element at its 5' end and a splice donor sequence 3'. This gene cassette is located in the LTRs of the retroviral vector. The other marker (neo) also 25 contains a promoter element but has a polyadenylation signal present at the 3' end of the coding sequence and is positioned between the viral LTRs. Both selectable markers contain an initiator ATG for proper translation. The design of VICTR 12 allows for the assessment of absolute titer as 30 assayed by the number of colonies resistant to antibiotic selection for the constitutively expressed marker possessing a polyadenylation signal. This titer can then be compared to that observed for gene-trapping and stable expression of the resistance marker flanked at its 3' end by a splice donor. 35 These numbers are important for the calculation of gene trapping frequency in the context of both nonspecific binding by retroviral integrase and directed binding by chimeric

integrase fusions. In addition, it provides an option to focus on the actual integration sites through infection and selection for the marker containing the polyadenylation signal. This eliminates the need for the fusion protein 5 binding to occur upstream and in the proximity of the target gene. Theoretically, any transcription factor binding sites present within the genome are targets for proximal integration and subsequent antibiotic resistance. Analysis of sequences flanking the LTRs of the retroviral vector 10 should reveal canonical factor binding sites. In addition, by including the promoter/splice donor design of VICTR 3, gene-trapping abilities are retained in VICTR 12.

VICTR A is a vector which does not contain gene trapping constructs but rather a selectable marker possessing all of the required entities for constitutive expression including, but not limited to, a promoter element capable of driving expression in eukaryotic cells and a polyadenylation and transcriptional terminal signal. Similar to VICTR 12, downstream gene trapping is not necessary for successful selection using VICTR A. This vector is intended solely to select for successful integrations and serves as a control for the identification of transcription factor binding sites flanking the integrant as mentioned above.

Finally, VICTR B is similar to VICTR A in that it
25 comprises a constitutively expressed selectable marker, but it also contains the bacterial β -lactamase ampicillin resistance selectable marker and a ColEl origin of replication. These entities allow for the rapid cloning of sequences flanking the long terminal repeats through
30 restriction digestion of genomic DNA from infected cells and ligation to form plasmid molecules which can be rescued by bacterial transformation, and subsequently sequenced. This vector allows for the rapid analysis of cellular sequences that contain putative binding sites for the transcription
35 factor of interest.

Other vector designs contemplated by the present invention are engineered to include an inducible regulatory

elements such as tetracycline, ecdysone, and other steroidresponsive promoters (No et al., Proc Natl Acad Sci USA
93:3345-51, 1996; Furth et al., Proc Natl Acad Sci USA
91:9302-6, 1994). These elements are operatively positioned
to allow the inducible control of expression of either the
selectable marker or endogenous genes proximal to site of
integration. Such inducibility provides a unique tool for
the regulation of target gene expression.

All of the gene trap vectors of the VICTR series, with 10 the exception of VICTRs A and B, are designed to form a fusion transcript between vector encoded sequence and the trapped target gene. All of the flanking exons may be sequenced according to the methods described in the following section. To facilitate sequencing, specific sequences are 15 engineered onto the ends of the selectable marker (e.g., puromycin coding region). Examples of such sequences include, but are not limited to unique sequences for priming PCR, and sequences complementary to standard M13 sequencing primers. Additionally, stop codons are added in all three 20 reading frames to ensure that no anomalous fusion proteins are produced. All of the unique 3' primer sequences are immediately followed by a synthetic 9 base pair splice donor sequence. This keeps the size of the exon comprising the selectable marker at a minimum to ensure proper splicing, and 25 positions the amplification and sequencing primers immediately adjacent to the flanking trapped exons to be sequenced as part of the generation of the collection of cells representing mutated transcription factor targets.

Since a cryptic splice donor sequence is found in the 30 inverted LTRs, this cryptic splice donor sequence has been removed from the VICTR vectors by site specific mutagenesis. It was deemed necessary to remove this splice donor so that it would not affect trapping associated splicing events.

When any members of the VICTR series are packaged into 35 infectious virus, the direction of transcription of the selectable marker is opposite to that of the direction of the normal transcription of the retrovirus. The reason for this

organization is that the regulatory elements such as the polyadenylation signal, the splice sites and the promoter elements found in the various members of the VICTR series can interfere with the transcription of the retroviral genome in 5 the packaging cell line. This potential interference may significantly reduce retroviral titers.

Although specific gene trapping vectors have been discussed at length above, the invention is by no means to be limited to such vectors. Several other types of vectors that 10 may also be used to incorporate relatively small engineered exons into a target cell transcripts include, but are not limited to, adenoviral vectors, adenoassociated virus vectors, SV40 based vectors, and papilloma virus vectors. Additionally, DNA vectors may be directly transferred into 15 the target cells using any of a variety of biochemical or physical means such as lipofection, chemical transfection, retrotransposition, electroporation, and the like.

Although, the use of specific selectable markers has been disclosed and discussed herein, the present invention is in no way limited to the specifically disclosed markers. Additional markers (and associated antibiotics) that are suitable for either positive or negative selection of eukaryotic cells are disclosed, inter alia, in Sambrook et al. (1989) Molecular Cloning Vols. I-III, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York, and Current Protocols in Molecular Biology (1989) John Wiley & Sons, all Vols. and periodic updates thereof, as well as Table I of U.S. Patent No. 5,464,764 issued November 7, 1995, the entirety of which is herein incorporated by reference. Any of the disclosed markers, as well as others known in the art, may be used to practice the present invention.

5.2. The Analysis of Mutated Genes and Transcripts

The presently described invention allows for large-scale 35 genetic analysis of the genomes of any organism for which there exists cultured cell lines. The Library may be constructed from any type of cell that can be transfected by

standard techniques or infected with recombinant retroviral vectors.

Where mouse ES cells are used, then the Library becomes a genetic tool able to completely represent mutations in 5 essentially every gene of the mouse genome. Since ES cells can be injected back into a blastocyst and become incorporated into normal development and ultimately the germ line, the cells of the Library effectively represent a complete panel of mutant transgenic mouse strains (see 10 generally, U.S. Patent No. 5,464,764 issued November 7, 1995, herein incorporated by reference).

A similar methodology may be used to construct virtually any non-human transgenic animal (or animal capable of being rendered transgenic). Such nonhuman transgenic animals may 15 include, for example, transgenic pigs, transgenic rats, transgenic rabbits, transgenic cattle, transgenic goats, and other transgenic animal species, particularly mammalian species, known in the art. Additionally, bovine, ovine, and porcine species, other members of the rodent family, e.g. 20 rat, as well as rabbit and guinea pig and non-human primates, such as chimpanzee, may be used to practice the present invention.

Transgenic animals produced using the presently described library and/or vectors are useful for the study of 25 basic biological processes and diseases including, but not limited to, aging, cancer, autoimmune disease, immune disorders, alopecia, glandular disorders, inflammatory disorders, diabetes, arthritis, high blood pressure, atherosclerosis, cardiovascular disease, pulmonary disease, 30 degenerative diseases of the neural or skeletal systems, Alzheimer's disease, Parkinson's disease, asthma, developmental disorders or abnormalities, infertility, epithelial ulcerations, and microbial pathogenesis (a relatively comprehensive review of such pathogens is provided, inter alia, in Mandell et al., 1990, "Principles and Practice of Infectious Disease" 3rd. ed., Churchill Livingstone Inc., New York, N.Y. 10036, herein incorporated

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by reference). As such, the described animals and cells are particularly useful for the practice of functional genomics.

5.2.1. Constructing a Library of Individually Mutated Cell Clones

were used to infect (or transfect) cells in culture, for example, mouse embryonic stem (ES) cells. Gene trap insertions were initially identified by antibiotic resistance (e.g., puromycin). Individual clones (colonies) were moved from a culture dish to individual wells of a multi-welled tissue culture plate (e.g. one with 96 wells). From this platform, the clones were be duplicated for storage and subsequent analysis. Each multi-well plate of clones was then processed by molecular biological techniques described in the following section in order to derive sequence of the gene that has been mutated. This entire process is presented schematically in Figure 4 (described below).

5.2.2. Identifying and Sequencing the Tagged Genes in the Library.

The relevant nucleic acid (and derived amino acid sequence information) will largely be obtained using PCR-based techniques that rely on knowing part of the sequence of the fusion transcripts (see generally, Frohman et al., 1988, Proc. Natl. Acad. Sci. U.S.A. 85(23):8998-9000, and U.S. Patents Nos. 4,683,195 to Saiki et al., and 4,683,202 to Mullis, which are herein incorporated by reference). Typically, such sequences are encoded by the foreign exon containing the selectable marker. The procedure is represented schematically in Figure 2 (3' RACE). Although each step of the procedure may be done manually, the procedure is also designed to be carried out using robots that can deliver reagents to multi well culture plates (e.g., but not limited to, 96-well plates).

The first step generates single stranded complementary DNA which is used in the PCR amplification reaction (Figure

The RNA substrate for cDNA synthesis may either be total cellular RNA or an mRNA fraction; preferably the latter. mRNA was isolated from cells directly in the wells of the tissue culture dish. The cells were lysed and mRNA was bound 5 by the complementary binding of the poly-adenylate tail to a poly-thymidine-associated solid matrix. The bound mRNA was washed several times and the reagents for the reverse transcription (RT) reaction were added. cDNA synthesis in the RT reaction was initiated at random positions along the 10 message by the binding of a random sequence primer (RS). This RS primer has approximately 6-9 random nucleotides at the 3' end to bind sites in the mRNA to prime cDNA synthesis, and a 5' tail sequence of known composition to act as an anchor for PCR amplification in the next step. 15 therefore no specificity for the trapped message in the RT step. Alternatively, a poly-dT primer appended with the specific sequences for the PCR may be used. Synthesis of the first strand of the cDNA initiates at the end of each trapped gene. At this point in the procedure, the bound mRNA may be 20 stored (at between about -70° C and about 4° C) and reused multiple times. Such storage is a valuable feature where one subsequently desires to analyze individual clones in more detail. The bound mRNA may also be used to clone the entire transcript using PCR-based protocols.

25 Specificity for the trapped, fusion transcript is introduced in the next step, PCR amplification. The primers for this reaction are complementary to the anchor sequence of the RS primer and to the selectable marker. Double stranded fragments between a fixed point in the selectable marker gene 30 and various points downstream in the appended transcript sequence are amplified. It is these fragments which will become the substrates for the sequencing reaction. The various end-points along the transcript sequence were determined by the binding of the random primer during the RT 35 reaction. These PCR products were diluted into the sequencing reaction mix, denatured and sequenced using a primer specific for the splice donor sequences of the gene

trap exon. Although, standard radioactively labeled nucleotides may be used in the sequencing reactions, sequences will typically be determined using standard dye terminator sequencing in conjunction with automated 5 sequencers (e.g., ABI sequencers and the like).

Several fragments of various sizes may serve as substrates for the sequencing reactions. This is not a problem since the sequencing reaction proceeds from a fixed point as defined by a specific primer sequence. Typically, 10 approximately 200 nucleotides of sequence were obtained for each trapped transcript. For the PCR fragments that are shorter than this, the sequencing reaction simply 'falls off' Sequences further 3' were then covered by the longer fragments amplified during PCR. One problem is 15 presented by the anchor sequences 'S' derived from the RS primer. When these are encountered during the sequencing of smaller fragments, they register as anomalous dye signals on the sequencing gels. To circumvent this potential problem, a restriction enzyme recognition site is included in the S 20 sequence. Digestion of the double stranded PCR products with this enzyme prior to sequencing eliminates the heterologous S sequences.

5.2.3. Identifying the Tagged Genes by Chromosomal Location

Any individually tagged gene may also be identified by PCR using chromosomal DNA as the template. To find an individual clone of interest in the Library arrayed as described above, genomic DNA is isolated from the pooled clones of ES cells as presented in Figure 3. One primer for the PCR is anchored in the gene trap vector, e.g., a puro exon-specific oligonucleotide. The other primer is located in the genomic DNA of interest. This genomic DNA primer may consist of either (1) DNA sequence that corresponds to the coding region of the gene of interest, or (2) DNA sequence from the locus of the gene of interest. In the first case, the only way that the two primers used may be juxtaposed to

give a positive PCR results (e.g., the correct size double-stranded DNA product) is if the gene trap vector has inserted into the gene of interest. Additionally, degenerate primers may be used, to identify and isolate related genes of interest. In the second case, the only way that the two primers used may be juxtaposed to provide the desired PCR result is if the gene trap vector has inserted into the region of interest that contains the primer for the known marker.

- 10 For example, if one wishes to obtain ES cell clones from the library that contain mutated genes located in a certain chromosomal position, PCR primers are designed that correspond to the *puro* gene (the puro-anchored primer) and a primer that corresponds to a marker known to be located in
- 15 the region of interest. Several different combinations of marker primers and primers that are located in the region of interest may also be used to obtain optimum results. In this manner, the mutated genes are identified by virtue of their location relative to sets of known markers. Genes in a
- 20 particular chromosomal region of interest could therefore be identified. The marker primers could also be designed correspond to sequences of known genes in order to screen for mutations in particular genes by PCR on genomic DNA templates. While this method is likely to be less
- 25 informative than the RT-PCR strategy described below, this technique would be useful as a alternative strategy to identify mutations in known genes. In addition, primers that correspond to sequence of known genes could be used in PCR reactions with marker-specific primers in order to identify
- 30 ES cell clones that contain mutations in genes proximal to the known genes. The sensitivity of detection is adequate to find such events when positive clones are subsequently identified as described below in the RT-PCR strategy.

35 5.3. A Sequence Database Identifies Genes Mutated in the Library.

Using the procedures described above, approximately 200

to about 600 bases of sequence from the cellular exons appended to the selectable marker exon (e.g., puro exon in VICTR vectors) may be identified. These sequences provide a means to identify and catalogue the genes mutated in each 5 clone of the Library. Such a database provides both an index for the presently disclosed libraries, and a resource for discovering novel genes. Alternatively, various comparisons can be made between the Library database sequences and any other sequence database as would be familiar to those 10 practiced in the art.

The novel utility of the Library lies in the ability for a person to search the Library database for a gene of interest based upon some knowledge of the nucleic acid or amino acid sequence. Once a sequence is identified, the 15 specific clone in the Library can be accessed and used to study gene function. This is accomplished by studying the effects of the mutation both in vitro and in vivo. For example, cell culture systems and animal models (i.e., transgenic animals) may be directly generated from the cells 20 found in the Library as will be familiar to those practiced in the art.

Additionally, the sequence information may be used to generate a highly specific probe for isolating both genomic clones from existing data bases, as well as a full length 25 cDNA. Additionally, the probe may be used to isolate the homologous gene from sufficiently related species, including humans. Once isolated, the gene may be over expressed, or used to generate a targeted knock-out vector that may be used to generate cells and animals that are homozygous for the 30 mutation of interest. Such animals and cells are deemed to be particularly useful as disease models (i.e., cancer, genetic abnormalities, AIDS, etc.), for developmental study, to assay for toxin susceptibility or the efficacy of therapeutic agents, and as hosts for gene delivery and 35 therapy experiments (e.g., experiments designed to correct a specific genetic defect in vivo).

5.4. Accessing Clones in the Library by a Pooling and Screening Procedure.

An alternative method of accessing individual clones is by searching the Library database for sequences in order to isolate a clone of interest from pools of library clones. The Library may be arrayed either as single clones, each with different insertions, or as sets of pooled clones. That is, as many clones as will represent insertions into essentially every gene in the genome are grown in sets of a defined number. For example, 100,000 clones can be arrayed in 2,000 sets of 50 clones. This can be accomplished by titrating the number of VICTR retroviral particles added to each well of 96-well tissue culture plates. Two thousand clones will fit on approximately 20 such plates. The number of clones may be dictated by the estimated number of genes in the genome of the cells being used. For example, there are approximately 100,000 genes in the genome of mouse ES cells. Therefore, a Library of mutations in essentially every gene in the mouse genome may be arrayed onto 20 96-well plates.

To find an individual clone of interest in the Library arrayed in this manner, reverse transcription-polymerase chain reactions (RT-PCR) are performed on mRNA isolated from pooled clones as presented in Figure 4. One primer for RT-PCR is anchored in the gene trap vector, i.e. a puro exonspecific oligonucleotide. The other primer is located in the cDNA sequence of a gene of interest. The only way that these two sequences can be juxtaposed to give a positive RT-PCR result (i.e. double stranded DNA fragment visible by agarose gel electrophoresis, as will be familiar to anyone practiced in the art) is by being present in a transcript from a gene trap event occurring in the gene of interest.

For example, if one wishes to obtain an ES cell clone with a mutation in the p53 gene, PCR primers are designed that correspond to the puro and p53 genes. If a VICTR trapping vector integrates into the p53 locus and results in the formation of a fusion mRNA, this mRNA may be detected by RT-PCR using these specifically designed primer pairs. The

sensitivity of detection is adequate to find such an event when positive cells are mixed with a large background of negative cells. The individual positive clones are subsequently identified by first locating the pool of 50 5 clones in which it resides. This process is described in Figure 5. The positive pool, once identified, is subsequently plated at limiting dilution (approximately 0.3 cells/well) such that individual clones may be isolated. To find the one positive event in 50 clones represented by this 10 pool, individual clones are isolated and arrayed on a 96-well plate. By pooling in columns and rows, the positive well containing the positive clone can be identified with relatively few RT-PCR reactions.

In addition to RT-PCR, the pools may be screened by 15 hybridization techniques (see generally Sambrook et al., 1989, Molecular Cloning: H Laboratory Manual 2nd edition, Cold Spring Harbor Press, Cold Spring Harbor, and Current Protocols in Molecular Biology, 1995, Ausubel et al. eds., John Wiley and Sons). Specific PCR fragments are generated 20 from the mutated genes essentially as described above for the sequencing protocols of the individual clones (first-strand synthesis using RT primed by a random or oligo dT primer that is appended to a specific primer binding site). trap DNA is amplified from the primer sets in the puro gene 25 and the specific sequences appended to the RT primer. this were done with pools, the resulting pooled set of amplified DNA fragments could be arrayed on membranes and probed by radioactive, or chemically or enzymatically labeled, hybridization probes specific for a gene of 30 interest. A positive radioactive result indicates that the gene of interest has been mutated in one of the clones of the positively-labeled pool. The individual positive clone is subsequently identified by PCR or hybridization essentially as outlined above.

Alternatively, a similar strategy may be used to identify the clone of interest from multiple plates, or any scheme where a two or three dimensional array (e.g., columns

and rows) of individual clones are pooled by row or by column. For example, 96 well plates of individual clones may be arranged adjacent to each other to provide a larger (or virtual/figurative) two dimensional grid (e.g., four plates 5 may be arranged to provide a net 16x24 grid), and the various rows and columns of the larger grid may be pooled to achieve substantially the same result.

Similarly, plates may simply be stacked, literally or figuratively, or arranged into a larger grid and stacked to 10 provide three dimensional arrays of individual clones. Representative pools from all three planes of the three dimensional grid may then be analyzed, and the three positive pools/planes may be aligned to identify the desired clone. For example, ten 96 well plates may be screened by pooling 15 the respective rows and columns from each plate (a total of 20 pools) as well as pooling all of the clones on each specific plate (10 additional pools). Using this method, one may effectively screen 960 clones by performing PCR on only 30 pooled samples.

The example provided below is merely illustrative of the subject invention. Given the level of skill in the art, one may be expected to modify any of the above or following disclosure to produce insubstantial differences from the specifically described features of the present invention. As such, the following example is provided solely by way of illustration and is not included for the purpose of limiting the invention in any way whatsoever.

6.0. EXAMPLES

30 6.1. Use of VICTR Series Vectors to Construct a Mouse ES cell Gene Trap Library

VICTR 3 was used to gather a set of gene trap clones. A plasmid containing the VICTR 3 cassette was constructed by conventional cloning techniques and designed to employ the features described above. Namely, the cassette contained a PGK promoter directing transcription of an exon that encodes the puro marker and ends in a canonical splice donor

sequence. At the end of the puromycin exon, sequences were added as described that allow for the annealing of two nested PCR and sequencing primers. The vector backbone was based on pBluescript KS+ from Stratagene Corporation.

The plasmid construct linearized by digestion with Sca I which cuts at a unique site in the plasmid backbone. The plasmid was then transfected into the mouse ES cell line AB2.2 by electroporation using a BioRad Genepulser apparatus. After the cells were allowed to recover, gene trap clones were selected by adding puromycin to the medium at a final concentration of 3 μ g/mL. Positive clones were allowed to grow under selection for approximately 10 days before being removed and cultured separately for storage and to determine the sequence of the disrupted gene.

Total RNA was isolated from an aliquot of cells from 15 each of 18 gene trap clones chosen for study. micrograms of this RNA was used in a first strand cDNA synthesis reaction using the "RS" primer. This primer has unique sequences (for subsequent PCR) on its 5' end and nine 20 random nucleotides or nine T (thymidine) residues on it's 3' Reaction products from the first strand synthesis were added directly to a PCR with outer primers specific for the engineered sequences of puromycin and the "RS" primer. After amplification, an aliquot of reaction products were subject 25 to a second round of amplification using primers internal, or nested, relative to the first set of PCR primers. second amplification provided more reaction product for sequencing and also provided increased specificity for the specifically gene trapped DNA.

30 The products of the nested PCR were visualized by agarose gel electrophoresis, and seventeen of the eighteen clones provided at least one band that was visible on the gel with ethidium bromide staining. Most gave only a single band which is an advantage in that a single band is generally 35 easier to sequence. The PCR products were sequenced directly after excess PCR primers and nucleotides were removed by filtration in a spin column (Centricon-100, Amicon). DNA was

added directly to dye terminator sequencing reactions
(purchased from ABI) using the standard M13 forward primer a
region for which was built into the end of the puro exon in
all of the PCR fragments. Thirteen of the seventeen clones
that gave a band after the PCR provided readable sequence.
The minimum number of readable nucleotides was 207 and some
of the clones provided over 500 nucleotides of useful
sequence.

Sample data from this set of clones is presented in

10 Figure 6. Only a portion of sequence (nucleotide or putative amino acid) for 9 Library clones obtained by the methods described in this invention are presented. Under each sequence fragment in the figure is aligned a homologous sequence that was identified using the BLAST (basic local alignment search tool) search algorithm (Altschul et al., 1990, J. Mol. Biol. 215:403-410).

In addition to known sequences, many new genes were also identified. Each of these sequences is labeled "OST" for "Omnibank Sequence Tags." OMNIBANK™ shall be the trademark 20 name for the Libraries generated using the disclosed technology.

These data demonstrate that the VICTR series vectors may efficiently trap genes, and that the procedures used to obtain sequence are reliable. With simple optimization of each step, it is presently possible to mutate every gene in a given population of cells, and obtain sequence from each of these mutated genes. The sample data provided in this example represents a small fraction of an entire Library. By simply performing the same procedures on a larger scale (with automation) a Library may be constructed that collectively comprises and indexes mutations in essentially every gene in the genome of the target cell.

Additional studies have used both VICTR 3 and VICTR 20.

Like VICTR 3, VICTR 20 is exemplary of a family of vectors

35 that incorporate two main functional units: a sequence acquisition component having a strong promoter element (phosphoglycerate kinase 1) active in ES cells that is fused

to the puromycin resistance gene coding sequence which lacks a polyadenylation sequence but is followed by a synthetic consensus splice donor sequence (PGKpuroSD); and 2) a mutagenic component that incorporates a splice acceptor 5 sequence fused to a selectable, colorimetric marker gene and followed by a polyadenylation sequence (for example, SAβgeopA or SAIRESβgeopA). Also like VICTR 3, stop codons have been engineered into all three reading frames in the region between the 3' end of the selectable marker and the splice 10 donor site. A diagrammatic description of structure and functions of VICTRs 3 and 20 is provided in Figure 7.

When VICTRs 3 and 20 were used in the commercial scale application of the presently disclosed invention, over 3,000 mutagenized ES cell clones were rapidly engineered and 15 obtained. Sequence analysis obtained from these clones has identified a wide variety of both previously identified and novel sequences. A representative sampling of previously known genes that were identified using the presently described methods is provided in Figure 8. The power of the 20 presently described invention as a genomics resource becomes apparent when one considers that the genes listed in Figure 8 were obtained and identified in less than a year whereas the references associated with the identification of the known genes span a period of roughly two decades. 25 importantly, the majority of the sequences thus far identified are novel, and, because of the functional aspects of the presently described ES cell system, the cellular and developmental functions of these novel sequences can be rapidly established.

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7.0. Reference to Microorganism Deposits

The following plasmids have been deposited at the American Type Culture Collection (ATCC), Rockville, MD, USA, under the terms of the Budapest Treaty on the International 35 Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and Regulations thereunder (Budapest Treaty) and are thus maintained and made available according

to the terms of the Budapest Treaty. Availability of such plasmids is not to be construed as a license to practice the invention in contravention of the rights granted under the authority of any government in accordance with its patent 5 laws.

The deposited cultures have been assigned the indicated ATCC deposit numbers:

	<u>Plasmid</u>	ATCC No.
	plex	97748
10	pExonII	97749
10	ppuro7	97750
	ppuro5	97751
	ppurol1	97752
	ppuro10	97753

All publications and patents mentioned in the above specification are herein incorporated by reference. Various modifications and variations of the described method and system of the invention will be apparent to those skilled in the art without departing from the scope and spirit of the invention. Although the invention has been described in connection with specific preferred embodiments, it should be understood that the invention as claimed should not be unduly limited to such specific embodiments. Indeed, various modifications of the above-described modes for carrying out the invention which are obvious to those skilled in the field of molecular biology or related fields are intended to be within the scope of the following claims.

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MICROORGANISMS
Optional Sheet in connection with the microorganism referred to on page 40, lines 5-25 of the description '
A. IDENTIFICATION OF DEPOSIT
Further deposits are identified on an additional sheet '
Name of depositary institution •
American Type Culture Collection
Address of depositary institution (including postal code and country) *
12301 Parklawn Drive
Rockville, MD 20852 US
Date of deposit ' October 9, 1996 Accession Number ' 97748
B. ADDITIONAL INDICATIONS (leave blank if not applicable). This information is continued on a separate attached sheet
C. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE " (if the indications are not all designated States)
D. SEPARATE FURNISHING OF INDICATIONS ((leave blank if not applicable)
The indications listed below will be submitted to the International Bureau later ' (Specify the general nature of the indications e.g., "Accession Number of Deposit")
E. This sheet was received with the International application when filed (to be checked by the receiving Office)
(Authorized Officer)
☐ The date of receipt (from the applicant) by the International Bureau *
was
(Authorized Officer)

Form PCT/RO/134 (January 1981)

International Application No: PCT/ /

Form PCT/RO/134 (cont.)

American Type Culture Collection

12301 Parklawn Drive Rockville, MD 20852 US

Accession No.	Date of Deposit
97749	October 9, 1996
97750	October 9, 1996
97751	October 9, 1996
97752	October 9, 1996
97753	October 9, 1996

CLAIMS

What is claimed is:

1. A library of cultured eucaryotic cells made by a process comprising the steps of:

- a) treating a first group of cells to stably integrate a first vector that mediates the splicing of a foreign exon internal to a cellular transcript;
- b) treating a second group of cells to stably integrate a second vector that mediates the splicing of a foreign exon
 10 5' to an exon of a cellular transcript; and
 - c) selecting for transduced cells that express the products encoded by the foreign exons.
- 2. A library according to claim 1 wherein said treating 15 is transfection.
 - 3. A library according to claim 1 wherein said treating is by infection.
- 4. A library according to claim 1 wherein said treating is by retrotransposition.
 - 5. A library according to any one of claims 1 through 4 wherein said cells are animal cells.

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- 6. A library according to claim 5 wherein said animal is mammalian.
- 7. A library according to claim 6 wherein said cells 30 are rodent cells.
 - 8. The use of a mutated cell from a library according to claim 6 to generate a non-human transgenic animal.
- 9. A vector for replacing the 3' end of an animal cell transcript with a foreign exon, comprising:
 - a) a selectable marker;

b) a splice acceptor site operatively positioned 5' to the initiation codon of said selectable marker;

- c) a polyadenylation site operatively positioned 3' to said selectable marker;
- 5 d) said vector not comprising a promoter element operatively positioned 5' of the coding region of said selectable marker; and
 - e) said vector not comprising a splice donor sequence operatively positioned between the 3' end of the coding region of said selectable marker and said polyadenylation site.
- 10. A vector for inserting foreign mutagenic
 polynucleotide sequence internal to animal cell transcripts,
 15 comprising:
 - a) a foreign exon;
 - b) a splice acceptor sequence operatively positioned 5' to the foreign exon;
 - c) a splice donor site operatively positioned 3' to said foreign exon;
 - d) a sequence comprising a nested set of stop codons in each of the three reading frames located between the 3' end of said foreign exon and said splice donor site;
- e) said vector not comprising a polyadenylation site operatively positioned 3' to said foreign exon; and
 - f) said vector not comprising a promoter element operatively positioned 5' to the coding region of said foreign exon.

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- 11. A vector for attaching a foreign exon upstream from the 3' end of an animal cell transcript, comprising:
 - a) a selectable marker;
 - b) a promoter element operatively positioned 5' to said selectable marker;
 - c) a splice donor site operatively positioned 3' to said selectable marker; and

d) said vector not comprising a transcription terminator or polyadenylation site operatively positioned relative to the coding region of said selectable marker; and

- e) said vector not comprising a splice acceptor site operatively positioned between said promoter element and the initiation codon of said selectable marker.
- 10 12. A vector according to claim 11 wherein said vector additionally comprises a foreign mutagenic polynucleotide sequence located upstream from said promoter.
- 13. A vector according to claim 12 wherein said vector 15 additionally comprises a splice acceptor operatively positioned upstream from said foreign mutagenic polynucleotide sequence.
- 14. A vector according to claim 13 wherein said foreign 20 mutagenic polynucleotide sequence comprises a polyadenylation site.
- 15. A vector according to claim 14, wherein said foreign mutagenic polynucleotide sequence additionally 25 comprises stop codons in all three reading frames.
- 16. A vector according to claim 12 in which a first recombinase recognition sequence is present upstream from said promoter and a second recombinase recognition sequence 30 is present downstream from said promoter.
 - 17. A vector according to any one of claims 9, 10, or 11 wherein said vector is a viral vector.
- 35 18. A vector according to claim 17 wherein said viral vector is a retroviral vector.

19. The use of a vector according to claim 9 to produce a library of mutated animal cells.

- 20. The use of a vector according to claim 10 to 5 produce mutated animal cells.
 - 21. The use of a vector according to claim 11 to produce mutated animal cells.
- 10 22. The use of a vector according to claim 11 to effect homologous recombination in an animal cell.
 - $\,$ 23. A stably transduced animal cell that incorporates a vector according to claim 16.

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- 24. A method of deleting a region of vector DNA from a cell according to claim 23, comprising:
 - a) providing a recombinase activity to the cell; and
- b) selecting for cells that lack the desired region of 20 vector DNA.
 - 25. A method of adding a region of DNA to a cell according to claim 23, comprising:
 - a) introducing the DNA to be added into the cell;
- a) providing a recombinase activity to the cell; and
 - b) selecting for cells that incorporate the added DNA.
 - 26. A method of effecting the inducible expression of a desired gene, comprising:
- 30 a) providing a cell according to claim 23 with a recombinase gene that is expressed by an inducible promoter; and
 - b) inducing said inducible promoter.
- 35 27. A method of gene discovery comprising:
 - a) adding a foreign polynucleotide to a population of target cells such that the foreign

polynucleotide is inserted throughout the genomes of the target cells; and

b) activating control elements encoded by the foreign polynucleotides that activate or repress the 5 expression of target cell genes that flank the integrated foreign polynucleotides, and identifying the regions of the target cell genome into which the foreign polynucleotides have integrated.

10 28. A library of cultured animal cells that stably integrate vectors according to claims 10 or 11.

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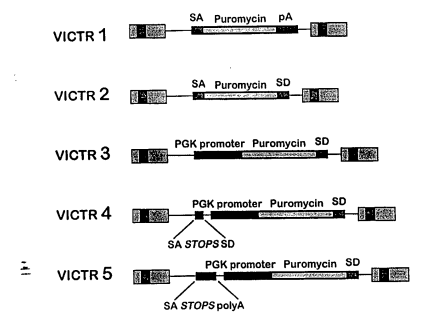


Figure 1

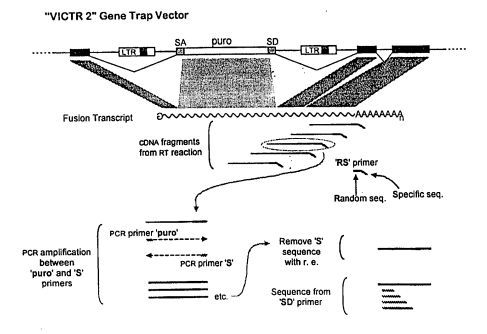


Figure 2

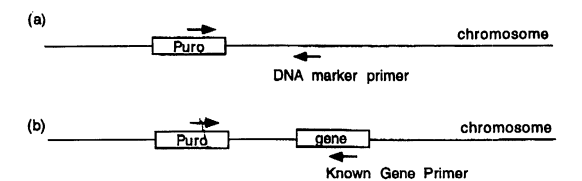


Figure 3

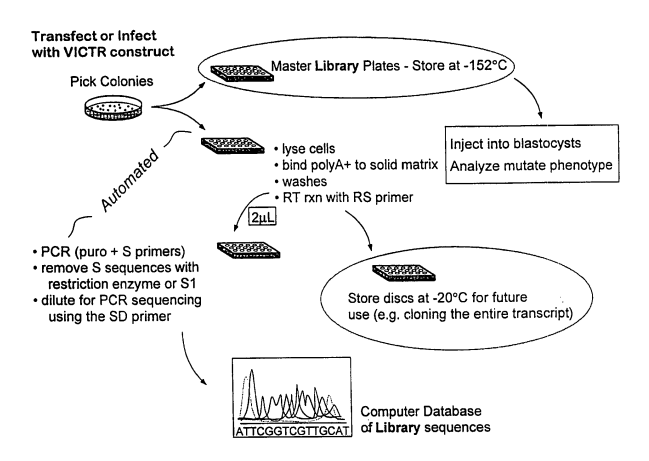
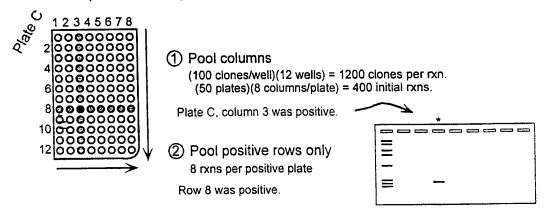


Figure 4

Identify Positive Pool

To screen all mouse genes (~100,000) with 5-fold redundancy would require about 50 plates of 96-wells (at 100 clones/well).



Identify Positive Clone

The pool on plate C, column 3, row 8 is thawed and plated as single clones:

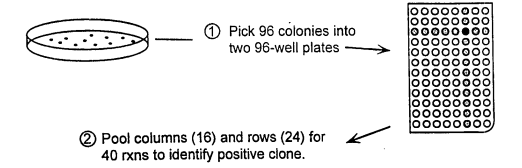


Figure 5

OST1:	248 TITATATAATATTTAATTTGTTTTACTGGGGTATATATGTGTGAAGAGGGCTTCT 302
rat GABA rho3:	1547 TITACATAATATITAATITGTTTTACTGGGGTATATATGTGTGAAGAGGACTTTT 1601
OST2:	56 ACCGTTGCGGAGGCTCACGTTTCTCAGATAGTACATCAGGTGTCATCGNTGTCAGAAGGT 115
mouse TCR-ATF1:	75 ACCGTTGCGGGGCCTCACGTTTCTCAGATAGTACATCAGGTGTCATCGTTATCAGAAAGT 134
OST3:	58 GIGMHHAGLHERDRKTVEELFXNCKVQVLIATSTLAWGVNFPAHLVIIKGTEYYDGKTRR 237 GIG+HHAGL ++DR +LF K+Q+LIATSTLAWGVN PAHLVIIKGT+++D K
Yeast ORF G9365:	1430 GIGLHHAGLVQKDRSISHQLFQKNKIQILIATSTLAWGVNLPAHLVIIKGTQFFDAKIEG 1489
OST4: seq. from US	137 GCGCAGAAGTGGTINCTGGAANTTTNTCCGCCNCCATCCAGTCTATTAATTGTTGACNGGA 196
patent 5470724:	166 GCGCAGAAGTGGTCCTGCAACTTTATCCGCCTCCATCCAGTCTATTAATTGTTGCCGGGA 225
OST5: mouse wnt-5A	108 TCWIRLGT*RXVGASLEYEYIRAS 179 TCW++L R VG +L+ +Y A+
protein precursor:	250 TCWLQLADFRKVGDALKEKYDSAA 273
OST6:	78 CTTATATGGCTACGGCGGCTTCAACATCTCCATTACACCCCAACTACAGCGTGTCCAGGCT 137
endopeptidase:	1407 CTTATATGGCTATGGCGGCTTCAACATATCCATCACACCCAACTACAGTGTTTCCAGGCT 1466
OST7:	109 AAAGCATGTAGCAGTTGTAGGACACACTAGACGAGAGCACCAGATCTCATTGTGGGTGG
mouse 45S pre rRNA:	1604 AAAGCATGTAGCAGTTGTAGGACACCAGAGCGAGGCACCAGATCTCATTGTGGGTGG
OST8:	161 TGGATGCAGNCTACCACTGTGTGGCTGCCCTATTTTACCTCAGTGCCTCAGTTCTGGAAG 220
rat MAL:	_
OST9:	103 ACCTGATTGTTATCCGTGCCTGCAGAAGTCCAGAAAATACAGACCAAAGTCAACCAGTA 162
mouse malic enzyme:	1666 ACCTGATTGTTATCCGTGGCCTGCAGAAGTCCAGAAAATACAGACCAAAGTCAACCAGTA 1725

Figure 6

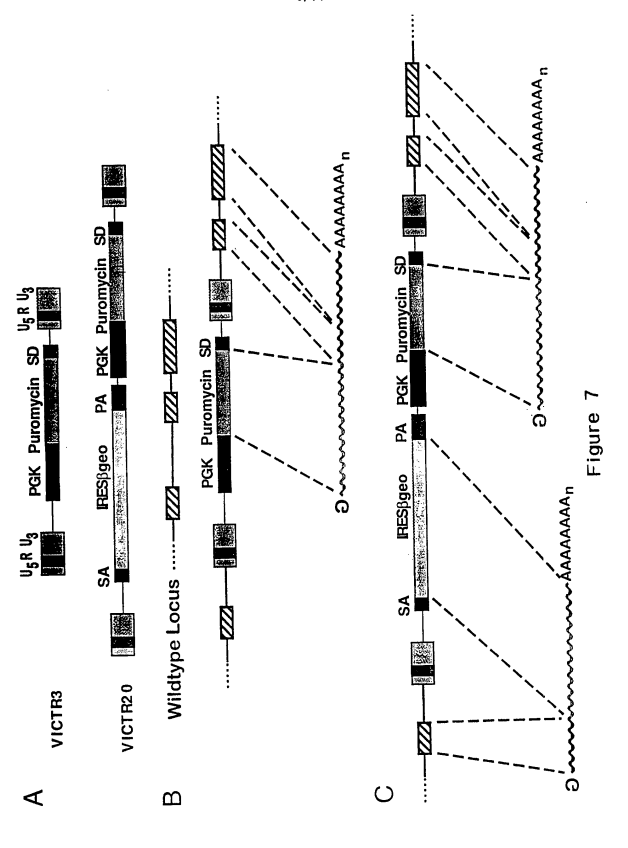


Figure 8

	Sequence Description	Mus musculus mu63102.rl Sonres mouse pJNHF19.5 Mus musculus CDAA clond 115.147 5.	Hus musculus House milit for retinal cyclic-GHP phosphodiesterase	Gamma-Subunic (CMP-PDE) (EC 5.1.4.17) Hus musculus Mouse mRNA	nus museculus aus museculus eriuo muso, Complete eds Aus museculus mj50b06.rl Soares mouse	embryo NDMEII.5 14.5 Mus musculus CDNA clone 479507 5.	Mus musculus Mouse mKNA for squalene synthase	Hus musculus M. musculus T. cell receptor alpha chain variable region	(v-aipha) Hus musculus mouse alpha-amylase-2	Nattus norvegitus Rat cytochrome P450	Mus musculus mg/7d10.rl Soares mouse embryo NbWEll.5 14.5 Mus musculus cDNA	Mus musculus House mouse; Musculus domesticus Postnatal (0 day) Brain mRNA for Ca2+ dependent activator	protein for secretion, complete cds Hus musculus mu5c11.rl Soares mouse lymph node NkHLM Hus musculus cDNA clone 641028 5' similar to TR:G294850	G294850 ALPHA-MUSCLE ACTIN Rattus norvegicus Rat TH-4 gene for	Mus musculus M.musculus 1gk-Vk2(70/3)	yene Hus musculus muttios,rl Soares mouse lymph node NDMLN Mus musculus cDNA	COURT OF SECULOR OF SECUEDAR OF SECULOR OF SECULOR OF SECULOR OF SECULOR OF SECULOR OF S	Ractus sp. EST110153 Ractus sp. CDNA	Homo sapiens 1151b07.s1 Soares pregnant ucture NableU Homo sapiens CDNA Lone 505429 J: similar to pric62249 6623249 CLEAVAGE pric62249 6623249 CLEAVAGE	Home sapiens Human mRNA for KIAA0242	Homo sapiens Human mRNA for KIAA0240	Home sapiens Human ser2 mRNA for RNA hinding profess CR2 complete one	Homo Sapiens 2106609.rl Stratagene Hela cell si 97/216 Nomo umpiens CDNA	Mus musculus mod9c0G.rl Life Tech mouse embryo 10 5dpc 10665016 Mus	musculus cDNA clone 556906 5' similar to gb.J05277 mouse hexokinase mRNA,	Entrus norvegicus for milla for ribonaria	Homo sapiens similar to glutamyl-tRHA contherant	Rattus sp. EST106973 Rattus sp. CDNA	Define Station to Symptomist The mouse embryo NDME11.5 14.5 Mus musculus CDNA	choine 442.09 5. Hus masculus Mouse 4.55 RNA gene Hus masculus mg74ell.rl Soares mouse embryo NKMEll 5 14.5 Mus musculus CDHA clone 418764 5.
	īđ.	196	306	834	986		305	156	101	621	966	199	17.6	168	156	88	196	881	976	841	921	831	734	B24		124	17	198	164	931
	pvalue	5.0e-133	2.6c 41	5.9e-48	1.9e-173		7.5e-71	3.0e-106	1.8c-70	4.06-34	1.4e-145	1.5e-45	2.6e-37	7.5e-112	1.06-126	1.7e-31	1.8e-178	7.3e-40	4.04-111	8.6e-154	2.0e-145	3.1e-161	1.2e-52	4.04-128		8.1c-143	4.8e-107	4.8c-38	1.84-81	1.2e~91 1.5e~141
- 1	UB Accession	96 W09445	9b r00746	90 088454	gb 028168 gb AA048968		95/022016	gb[x53732	95/3001/96	951CtH d2	gb AA003309	yb 086214	yb}AA189233	95 y y 00169	95 272384	gb AA190122	gb AA104745	ցելույյցու	gb AA156426	90 087684	45 [D87077	gb 028482	gb[AA114106	(PRIOIPY)rif		000508196	gb[C06148	gb 1132346	gb[AA009152	gb M12658 gb AA058245
	Onusilkank	osr4	os r 5	05T22	OST25		uST36	02F38	0ST41	0ST42	0ST45	05751	05156	OST74	OST75	05786	05795	96150	057117	OST118	0Sr119	0ST121	05:11))	osT154		971730	051193	051243	OST246	05726B
	The following table includes 586 OSTs. OSTs with hit into prodom and Genuenk	parented befores have been removed or west as sequence with repressive																												

057281	gb 065313	1.86-180	186	Mus musculus Mus musculus reserrivating 5813-domain	021562	y Ju
1000	004 004 014	4) June 60	2	cds according milytopl of Course manner	OSTSGB	4)q6
567197	90 1000 100	5		clone 477500 5: similar to go: 302809	0.5T-57.1	ماطو
				Mouse neural specific Calmodulin-binding protein P-57 MMMA,		
OST297	gb Z77585	3.00-168	101	Mus musculus A.musculus milità for thioredoxio	0.511572	₹ q6
021300	9b M75122	1.8e-203	98.5	Mus musculus Mouse acid beta- calactosidas (GLB-1) quie, exon 16	057573	gbit
021303	gb ₩34850	2.7e-97	116	Mustaction of the second of th		,
				clone 153067 5 similar to gh: Ull248 Mus musculus C578L/6J ribosomal	054577	op iv
0.5T411	gb w80427	3.0e-73	1.5R	protein 528 mkNA, complete Nomo sapiens zd82d06.sl Soares tetut	057581	a) dy
				heart NbHH19W Homo sapiens CDNA clond	051'582	0 96
os T 314	gb(T14710	4.0c-54	7.13	metastasis-associated protein mia-1 Hono appiens EST31842 Hono sapiens	(P. STEPS)	1
OST316	gb w11499	1.24-72	166	Aus musculus maddhús.rl Soares mouse	· ·	2
				PARMILES FOR MECELLE CON CLOSE PARMILE FOR SHICKER DOVIN	051593	11 96
05T3Z8	9b{W10861	3.74-59	168	REDUCTASE 7.2 KD PROTEIN WAS BUSSELUS BASSCOB.rl Scares House DINNELS. Uns massculus CDNA clone	057594	x 76
05T331	99101010	6.8e-119	831	314596 5. Mus musculus Mus musculus ablphilin-l	OSTS95	796
05T342	95/010120	3.1e-143	156	(abi-1) mRNA, complete cd:: Mus musculus Mus musculus SKD2 mRIA,	OCTS98	x)qG
0ST356	gb H60456	1.8e-117	924	complete cds Mus musculus Mouse cyclophilin mRHA,	051600	7
UST361	09ELLM qB	5.7e-37	306	complete cds Nus musculus me65fll.rl Scares mouse ambros Almantl 4 14 6 Mis musculus c1844	OST607	4 76
1100	Catal Indian	7 '41-184	47.6	Clone 400461 5. Mrs margular House mouse: Musculus	OSTÉLI	4107
99 (11.7)	700/00/00		;	clomestricus min for 14-3-1 tau.	-	,
051386	9668796	2.6e-35	85 E	Hustolius M.musculus 94kb genomic semimore encoding Tax onto	OSTG18	100
02F389	gb T51727	1.8e-78	89.8	Home sapiens yb28c11.11 Home sapiens	06,9450	4
OST401	95162M}96	J. Le-33	11.6	CDNA CIONE 72300 3 Mus mosculus malfacoll.rl Sources mouse nAnnyll S. Aus musculus CDNA clone		5
				148998 5' SIMITAE CO SWIYEFA_ECOLI P32054 CDF-D-HAMBOSE DEBYDRATASE		
057411	90/148542	2.00-68	78%	Homo sapiens yy49407.rl Homo sapiens	OST623	<u> </u>
UST418	gb 621163	1.7e-84	158	Homo sapiens human STS WI-15024	057626	Hdis
05T425	gb x04480	8.1c-58		Hus ausculus Mouse milita for	0007663	. 1
OSF4 30	T£678W dp	5.7c-93	196	Hus mustellus mc73407, r1 Soures mouse umbryo NbMK13,5 14.5 Mus museulus cPNA		
				cione 401244 5' similar to gb:N23419 Initiation Factor 5A (HUMAH)	;	
0.57439	gb M26756	2.46-134	878	Mus musculus Mouse malic enzyme mRUA. complete cds	057664	25
057442	gb[W2593B	2.60-49	701	Homo sapiens 15b8 Ihman recina culla randomly primed sublibrary Homo	057671	2100
OST448	65270Y dg	4.34-72	198	sapiens CDNA Homo sapiens H. sapiens mkNA for		-
0:-T5 31	ub[x95591	3.1e-206	41.4	Physical protects Mus musculus M. musculus mette for C10		<u>.</u>
051536	yb w75435	4.6e-75	156	NUS musculus meSOdO6.rl Souces mouse Nus musculus meSOdO6.rl Souces mouse embryo NUNED1.S 14.5 Mus musculus CDUA embryo 200001.cl = 1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1	OSTCB0 OST702	33
				Hus musculus chaperonin 10 mkHA. Complete cds (MOUSE)	051707	a qu
27 P. C.	10 141 140 11	6.8c 216	3.65	Mus musculus mb32g03.rl Source manner piNMF19.5 Nus musculus città ciente 311166 e cimilar es cuerenda veate	057716	4

Mus musculus M.musculus mRNA for sodium/botassium ATPase beta subunit	Nus musculus mg64a07.rl Soares mouse embryo NbHE13.5 14.5 Mus musculus CD4A	Grams anschlus mojiltoz.ri Life Tech mouse embryo lo Sdyc 10655016 Mus	musculus cuts atone 3/10/10/10/10/10/10/10/10/10/10/10/10/10/	CUMA Cione sobasou J Strus norvegicus Rattus norvegicus RNA polymerase II transcription factor	ALI DA GUOUTIC MARA, COMPLETE CUA MAR MUSCULUS MAÑA DOÑ. TI SOBRES MOUSE PLACETTE ATRANTO S. 14.5 MUS MUSCULUS PROVINCATOR ATRAIN S. 14.5 MUS MUSCULUS	Homo sapiens yqqquol2.rl Homo sapiens	Ratus ratus Rat manA for water Channel aquaporin 3 (AQP3), complete	Hus musculus Mus musculus domusticus coiled-coil protein (CG-1) mkNA.	complete cush meddad2.rl Soares mouse mbrow NUMEII.5 14.5 Mus musculus CDNA close 190114 5.	Mus musculus M.musculus mRNA for olycogen synthase	Rattus norvegicus Rattus norvegicus PSD-95/SAP90-associated protein-1	BRNA, COMPIECE COS Hus musculus Nouse BRNA for	Hos musculus Aus musculus histone Hos a (40% I) mans complete cds	Huss musculus ma35d03.rl Life Tech mouse brain Mus musculus CDNA clone 315673.5	Mus musculus mudbios.rl Soures mouse lymph node Nimilat Hus musculus clan. clone 642393 5: similar to gb:L00993	Hus musculus autoantigen La (MOUSE) Homo sapitens ymllf07.rl Homo sapitens	CDNA CIONE 4/32-5 5 Hust Buckdington Mus musculus mouse embryonic region Mus masculus	CURA clone 518938 5' cimilar to gp.L15599 Mus mesculus 'Y-box binding proceding Mid' 3' end (MOUSE)	Homo supiens th81b02.rl Soares fetal liver spleen lNFLS SI Homo supiens rous alone 477851 c.	Homo capiens arious mona capiens clone N9S Rep-8	Hus mesculus mil9u06.rl Soares mouse plWHF19.5 Hus musculus CDMA clone 461954 5: similar to gui-MIB775 House tau microcubule binding protein muMA.	complete (MOUSE) Mus musculus Mus musculus C57UL/63	Sec61 protein complex gamma subunit mRNA. complete cds Mus so NonGenon-MRN domain-containing	octumer-binding protein (mice, u-cell leukemia, BCLL, mRNA, 241 ncl. Muse mangel and 24,010 cl. States	plumily's Hus musculus contactore 130365 5.	Mus musculus Mouse stefnanto gene Homo sapiens ESTD1041 Homo sapiens CDNA clone HHCPB14 similar to	CAMP-regulated phosphoprotein Homo sapiens ym4ddl.rl Homo sapiens	cons crous males of the mouse musculus males mouse embryo NUMELLS 14.5 Mus musculus clita close 175304 5*
3.26	11.9	126	158	151	924	808	911	158	984	118	86¥	186	196	158	11.6	1.98	831		B1.	818	9 . 69 8	97.4	ž			95% 85%	821	196
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				Company of the comment of the commen	_				adhesion molecule 1
0511096	50 D87077	7.7e-112	88.	(CX31.1) indit, complete cos flow sapiens aman mRNA for KIAA0240	05T1376	gb x63615	1.1e-33	928	Mus musculus H. musculus Camk-2 mRNA for Ca2-Calmodulin dependent protein
5011350	gb w44423	1.00-66	198	gene, partial COS Homo sapiens 2028f04.sl Soutes Euroscent fibroblasts NMISF Home	OST1387	46(51552)	16.56.57	931	kinase II beta subunit Homo sapiens Human fetal beain cittle
2	94503×1540	311	, c	sapiens con clone 133647 3. Her marculus marculus scribb	0511410	C0965M qh	4.84.77	. G 8	5'-end GEN-149C01 Mus musculus md7]aU6.rl Soures mouse
0SF1137	gb #63485	1.26-89	946	Rattus norvegicus Rat matrin 3 mKHA Mus musculus Hat matrin 3 mKHA		-			embryo NbME13.5 14.5 Mus musculus cDNA clone 174002 5.
25	0.660M 95	6.4e-109	101	Hus musculus me90d10.rl Goares mouse enbryo NuMe13.5 145 Aug supscellus culA	0571419	gb{W10923	1.2e-122	11.6	Mus musculus ma40all.rl Soares mouse p3NMF19.5 Mus musculus cDNA clone
0211125	91-[101.0124	2. Je65	176	Clone 402815 S. SERRIAR TO 90:577531 INTERFERON-THUMCHBLE PROTEIN 1-80 Mus monscilus ma42dll.rl Stories monice	00301433	ญปุ่งหยุยชย	1,76-145	94%	Mus musculus m(77f07.rl Soares mouse embryo NiMell, 5 14.5 Mus musculus etrila ellono 420125.5
				DINGES State Busculus CDMA CAULUS DINGES STATE TO THE COURT OF T	05T1453	95/018312	5.60-95	199	Committee of the contraction of
05T116S	95 454649	1.9e-184	116	APOLIPOPROTEIN C-11 PRECURSON Hus musculus md07b12.rl Soares mouse embryo NDHE13.5 14.5 Mus musculus cD8A	0571457	90 878851	5.94-45	808	Common Septens yighted in Homo sapiens CDNA clone 146525 5
osr1179	9068004/96	8.54-84	944	clone 167679 5. Hus musculation mg99e02 rl Soares mouse	0511470	gb J05504	1.5e-136	954	Mus musculus Mouse guanylate cyclase/artial natriurecic factor RNA, complete cds
OST1186	gb U89840	2.8e-70	1.96	embryo NbMil.) 14.5 Mus musculus cuna Conne 441146 5: Mus musculus Mus musculus prograte corrector orderin (Prip44) mRNA.	OST1478	gb[U57821	B.4e-55	109	Mus musculus Hus musculus inhibitor of HyoD family-b (I-mf) makka, complete eds
osr1192	gb w82490	5.3127	196	complete cds Motorer 1500 memory Parent Pare	0571488	gb 024681	4.3e-235	176	synthetic construct Synthetic NADH:cytochrome c reductase fusion protein mRUA, complete cds
0ST1207	90 04063763	1.34-56	198	404094 5. Mus musculus mj79d10.rl Scares mouse o3NNF19-5 Hus musculus CDNA close	OST11492	gb W10703	3.6e-109	986	Mus musculus ma51gU5.rl Soares mouse plumF19.5 Mus musculus cDNA clone l14456 5'
0ST1223	gb AA002931	1.5e-189	166	482323 5' Mus mg40f08.rl Soares mouse embryo (MbMI).5 14.5 Mus musculus cDRA	0511493	gb[AA097483	3.54-197	186	Mus musculus mkl7d04.rl Soares mouse p3NNF19.5 Mus musculus CDRA clone 491159 5. similar to gb:L05093 60s R180SOMAL PROTEIN L18A (HUMAN)
05T1226	gb U37353	7.5e-279	958	Aus susculus Aus susculus protein	OST1499	gb T51184	1.34-60	941	Homo sapiens yb94h05.rl Homo sapiens cDNA clone 78873 5
0511233	101 W98701	6.36-91	988	prospinates of being marchal constructly subunit markh, parchal cds Mus musculus mg12f05.rl Soares mouse	0571504	gb L34260	1.7e-196	914	Mus musculus Mus musculus integral membrane protein 1 (Itml) mRNA,
	•			embryo NbME13.5 14.5 Mus musculus cDNA clone 423585 5'	05/11508	90103306	1.8e-164	941	complete cos Mus musculus Mus musculus core-binding factor mNN sections
OST1234	gb D85904	1.1e-180		Mus musculus Mouse mann lor apy-1,	OSF1520	yb W18420	4.7e-37	881	Hus musculus mb68e07.rl Soares mouse blanklight Aus musculus con clone
0ST1241	058600 198	4.66-184		(ZNF143) minnA, complete cds					334596 5' similar to PIR:JH0457 JH0457
0511247	gb AA051266	4.7e-126	9.76	Mus musculus mydinoz.rl Soares moose embryo NDMEI3.5 145 Mus musculus CDAA elone 478899 5' similar to SW:LlOK.RAT	OST1523	gb U23769	7.2e-195	921	Rattus norvegicus Rattus norvegicus CLP36 (clp36) mRNA, complete cds
osr1265	£7698X dg	6.80-183	306	Q05310 LEYDIG CELL TUMOR 10 KD PROTEIN Mus musculus M.musculus wRNA for	OST1554	gb w85270	2.3e-168	196	Hus musculus mf42d05.rl Soares mouse embryo NDHELL35 14.5 Hus musculus CDHA
0571267	90 184271	1.6e-32	198	Laup-l procesn Lamo sapienu yq22g02.rl Soares retina N2b4HH Humo sapienu CDNA clone 274730			:		SW.) PYK_BOVIN PJ7980 FROMGANIC PYKOPHOSHHATASE
;		ž ,	;	S' similar to SP:YPK1_YEAST P12688 SERINE/THREONNE-PROTEIN KINASE YPK1	0571556	gb AA117514	9.7e-150	e S	Mus musculus mn29c09.tl beddington mouse embryonic region Mus musculus CDNA clone 519344 5
OST1269	96[019977	4.06-130	2) 4	Homo Sapiens numan preparocarboxypuptidase A2 (proCPA2)	0571558	gb #33765	2.2c-67	198	Rattus sp. EST110085 Rattus sp. cDNA 5' end
osr1274	gb[H14634	2.1e-139	85.	minn, conveyious Rat mitochondrial propionyl-CoA carboxyluse (FCCase)	0571567	96)014636	1.74-39	941	Hus musculus Mus ausculus serine/lhreonine protein kinase DLK musca pomoleia ple
05T1294	£85£0£ 46	6.80-69	156	beta-subunit mkNA, complete cus Rattus norvegicus Hat clathrin heavy chain mkNA, complete cds	0571601	gh WJ9611	1.3e-89	851	Homo, sapiens zelyed8.rl Soares parathyroid tumor WhiPA Homo sapiens
05F1339	95 H97190	J.1e-118	834	Homo sapiens Human Sp2 protein mRRA,	0571603	ctb W09922	4.00-109	198	cDNA clone 322790 5. Mus musculus ma67e08.rl Soares mouse
05T1341	gb w89611	7.8e-142	933	Hus mocculus mf72h02.rl Soures mouse embryo NNFELL'S 14.5 Mos mouselvs child		-			plumf19.5 Mus musculus cDNA clone 115782 5's similar to PIR:244566 544566 Noothetical protein L8167.23 - yeast
osrt1354	gb W)4481	2.20-64	851	CLONE 412022 OF TABLE SE SOARES LEGAL HOME SAPERS 2475512 SE SOARES CHIA CLONE SAPERS CHIA CLONE	0571628	96/1051239	7.9e-78	166	Hus musculus Hus musculus lysosomal-ausociated multir fansmembrane protein (CAPTM)
05T1359	gh M65672	5.1e-35	3.6	Have musculus melliduy.rl Soures mouse embryo NbWEL).S 14.5 Was musculus cum clone 10717 5: similar to	0571649	06061904196	7.9e-158	196	mANA, complete cds Hus musculus mj46e07.rl Soares mouse embryo NDME11.5 14.5 Mus musculus cDNA
05T1369	955294	1.1c-109	17.6	SW:UBCS_UNDARE_PISTS UBJQUITH-CONJUCKTING ENTYME E2-17 KD UBJQUITH-CONJUCKTING ENTYME E2-17 KD	0571653	90 90 90	5.1e-143	1/6	Clone 479172 5. Hus musculus mi27f12.rl Soares mouse embryo NoMILL)5 14.5 Hus musculus CDHA
}				embryo NLME13.5 14.5 Mus musculus CDIA clone 374060 5' similar to QD:L06019	0ST1658	gb[w12941	5.2e-95	176	clone 464783 5: Was musculus ma89d07.rl Scares mouse

6.5c-138
4.7e-118 96%
2.40-66 80%
1.8e-35 85N
5.2c-109 9HV
3.8e-40 791
1.7e-144 938
1.1c-67 931
1.94-122 861
1.4e-55 93%
2.3e-86 97%
5.4e-61 951
3.1e-33 69%
4.1e-112 971
6.0u-153 93%
2.8e-109 94%
2.4e-9h 961
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1.26-4) 9.
8.3e-85 96t

SM:YA9E_SCHPO Q09790 HYPOTHETICAL 23.7 KD PHOTEN C13GE.14 IN CHROMOSOME I. Home Apriens H. sapiens partial CDNA	sequence; clone c-2gb07 Mus musculus Mouse F52 mkNA for a	novel protein Homo capiens x1911nd.rl Stratugen colon (#917204) Homo sapiens cDMA	Cione 512023 S. Rattus sp. Rat myoadenylate deaminase	(ANY Occupiests) many, complete cou New musculus Mouse mills for country of a complete city	Ayingtockijish J. Compacte Cos Akas musculus Mouse mRNA for HBp15/L22.	complete cas long sapiens yd04g12.rl Howo sapiens	CON CIONE TACE TO THE STATE OF THE PROPERTY OF	149116 5. Mus musculus mcOddO4.rl Soares mouse plNMF19.5 Mus musculus CDNA clone	Jaive, 3-1947 S. norvegicus mRNA for carnitine/acylcarnitine carrier	procein procenta dibbata mb24c06.rl Soares mouse placenta (dibbata), 5 14.5 Mus musculus	CDNA clone 44444 5 ilomo sapiens splicing	FACEOF (CCI.4) MKNA, COMPLETE CUS HUSS MUSICALUS MESCHIOLT SCATES MOUSE embroo NDME11.5 14.5 HUS MUSICALUS CDMA	Clone 418481 5' Mus musculus mj98h06.rl Soares mouse pjNMF19.5 Mus musculus CDNA clone	48418/ 5' 48418/	clone 440710 5' similar to gb:H31690 Mouse argininosuccinate synthetase Mouse argininosuccinate synthetase Mouse argininosuccinate synthetase		e embryor	HUS BUSCULUS mayBe09.rl Soares mouse plantPlo.5 Hus musculus cDRA clone plantPlo.5.	Nus musculus mj13d01.rl Soares mouse embryo NUME13.5 14.5 Mus musculus CDRA clone 475809 5 similar to cu. vac crumo nj1016 uvermiratis 1947	Mys musculus mb18b08.rl Soares mouse	plumF19.5 Mus musculus cDNA clone 111671 S.	CDNA clone 262491 S	Mus marculus mosfeol; r. l. life Tecti mouse embryo 8 5dpc 10664019 fus marculus cDNA clone 557595; s. similur to gb:N31642 WYPOXANTHIME-GUNNINE pticyptiopSyltayNSTERASE (UtbXNI); gb:J00423 mouse hypoxanthine	phosphoribosyltransferase (hprt) arna Nus musculus mardigod, rl Suarus mouse Albus musculus choric cons	148514 mascains mj98057.rl Shares mense panetals, 5 Hus mascalou (CHIA Colone	CE0514) COLLED COLL DOHATIS HORD SAPIEST SQUAGE HUMAN (\$3723) HORD SAPIEST COLL	clone 611978 5. Mus musculus Mus musculus co-chaperonin 'cofactor A' mRNA.	
198		781	924	941	858	85%	166	156	851	778	914	106	196	924	5			86	116	941			1.6	116	34.	106	196	15
7 8 34	7.00-66	5.10-85	1.2e-81	1.54.95	2.54-101	7.46-73	4.10-133	7.5e-58	1.9c-104	6.1e-J2	1.2e-105	9.50-88	8.3e-126	5.le-126	6	800.00	0.1	4.2e-122	3.7e-83	3.16-54	5	1. 16-4.	2.4e-164	1.76-55	1.00.58	4.2e-62	3.40-119	1.8e-116
ri ccoaldo	46813XJdg	gb AA100747	95 302811	ար Կողագրեր	gb D17653	76008T dp	990064 46	gb w18873	11878x dg	91/1/013837	gb {L10911	190\#87091	060080VV]05	gb AA015380	, 1350ml 42	95 5533408	60077199 05	gb w34469	gb AA049859	95 W14179		50 [425844	gb AA104747	UD[W35819	96]55061741	90 10167801	666500/46	gb w85263
1966430	OST2307	0512323	0ST2322	0072346	OST2347	05T2353	0512357	OST2361	0572367	0572368	0572379	0572380	0512381	OST2382	o c	300000	100	OST2400	0ST2401	0272416		0572418	05T2433	051/2442	0572447	0512455	0ST2459	UST:2464

0572829	90(2002649	7.7e-90	941	LPS-binding protein Hus musculus mg/8006 r1 Source mouse Authoris 6 14 6 Pure maneral in CP14					embryo NbME13.5 14.5 Mus musculus cbNA clone 479149 5' similar to WP:F45E12.4 CE02740
				CLORE 426106 5' SIRILAR CO SWINSES, ULUMAN QOOSB7 SERUM PROFEIU MCES, [1]	0512963	gb]W04744	4.2e-31	108	Homo sapiens 22/9c08.rl Soares fetal lung NbHL19W Homo sapiens cDNA clone 298766 5
OST2834	95/05/09	1.4e-222	17.6	Appropriate and other process of the control of the	0572971	gb[AA120487	9.2e-107	101	Mus musculus mul2f07.rl Beddington mouse embryonic region Mus musculus cDNA clone 537733 5° similar to
osrzaus	36 AA 0 6 0 7 9 5	2.1e-89	974	Mus musculus mj79405.1 Goarde mouse planyf19,5 Mus musculus CDM, close 48211,5 S. saminer to MP:F42H10.4		19.50 cm f. c.	601		SW:YDNS_YEAST F38219 HYPOTHETICAL 44 KD PROTEIN IN SCOZ-MKF1 INTERGENIC RECION OF THE PART
osasaso	96 AA163971	6.00-61	707	EZOUDO CHIL HVS masculus ms40a01.rl Life Tech mouse embryo 13 5dpc 10666014 Mus					neuroglycan C precursor aRNA, complete
0572842	9b #54515	6.10-64	911	musculus catha cione L11942. Hus masculus addbalo.tl Scarca mouse embryo NUME13.5 14.5 Hus musculus CDMA clone 18786 65 's shallar to dp.100718. APP-RIBSYLATION FATTOR-LIKE PROFEIN 3	02129	gb AA206420	1.26-71	85	Homo appiens registral Stratagene neuroepithelium (4937231) Nomo sapiens and companient (4937231) Nomo sapiens result close (49399) similar to TR: (49372006 6937006 MRAA: EXPRESED
OST2877	06 J03583	1.le-66	938	Rattus norvegicus Rat Clathrin heavy chain mRNA complete cds	0512983	gb[W49206	1.8c-119	186	SEQUENCE TAG Mus musculus mc91g12.rl Soares mous
OST2883	95 W34850	4.8e-75	931	Hus musculus mc62b02.rl Soares mouse embryo NDME13.5 14.5 Mus musculus CDNA clone 133067.5 similar to db:u11248 Mus musculus C57MLKA1 ribasomal					embryo NUMEll.5 14.5 Mus musculus CDNA clone 355942 S similar to PIR:S44900 S44900 ZK652.10 protein - Caenorhabditis elegans
0572892	95/464 96	1.40-125	186	protein S28 mRNA, complete (MOUSE) Mus musculus mg01e10 t.1 Soares acoust embrox NPMP11 5 14 5 Mus musculus cilik	0512987	gb AA027683	2.3e-134	196	Mus musculus mil2b01.rl Soares mouse plumF19.5 Mus musculus cDNA clone 463273 5'
				clone 422538 5' similar to gb:J04823_rnal CyTOCHROME C OXIDASE	0572988		2.24-52		Mus musculus domesticus Mouse testis-specific mRNA pBs6.2
UST2897	9b w11047	7.9e-132	978	POLYPEPTIDE VIII-LIVER/HEART Hus musculus ma78d10.rl Scares mouse planF19:5 Hus musculus cDNA clone 118819 5'	OST2989	95 AA152050	1.36-46	# 9/	Homo sapiens zisbiz.ri soares pregnant uterus NbHPU Homo sapiens CDNA clone 50\$151 5: similar to gb:H90156_cds1 TRANSCRIFTION FACTOR
OST2909	gb AA166258	B.94-120	196	Mus musculus ma49c09.rl Life Tuch mouse embryo 13 5dpc 1066014 Mus musculus CDNA clane 64896 5	OST2991	171E00AA)dy	8.4e-151	166	BTF3 (HUMAN) Mus musculus mg56h09.rl Soares mouse embryo NDME13.5 14.5 Mus musculus ci
0:r2911	gb U73478	1,46-117	864	Hus musculus Hus musculus acidic inclear phosphoprotein pp 2 mun.					clone 437057 5' similar to gb:#24194 GUANINE NUCLEOTIDE-BINDING PROTEIN BETA SUBLINIT-LIKE PROTEIN (HUMAN):
OST2914	96/012236	4.0e-136	156	Compare too Mus musculus Mus musculus AKR alpha M290 integrin akkA, complete cus	05T2994	96 851546	1.94-51	831	gb:X75313 m.muscutus Homo sapiens yg72h12.rl Homo sapiem
OST2916	gb D77002	1.4e-67	928	Mus musculus Mouse embryonal carcinoma F9 cell cDNA, 93E10					cDNA clone 38905 5' similar to SP:VILI_CHICK P02640
0572921	gb W75740	B.4c-106	986	Mus musculus me55b06.rl Soares mouse embryo NDME13.5 14.5 Mus musculus cDNA	0ST2996	gb x99921	1.6e-82		Mus musculus M.musculus mRNA for S100 calcium-binding protein All
0512922	gb 050544	8.4e-135	881	clone 191179 5' Homo sapiens Human lymphocyte mRNA for	0512998	gb[b19012	3.2e-48	104	Mus musculus Mouse 1'-directed CDNA, MUSCS01209, clone mc015
UST2923	gb w85631	3.2e-108	17.6	TFIID subunic p21. complete cds Mus musculus mf37b01.rl Soares mouse embryo NbME13.5 14.5 Mus musculus CDNA	505150	205/20 ap	1.36-109		nus musculus nus musculus iens major intriusic protein (MIP) munA, complete cds
OST2926	gb w59561	6.3e-164	273	clone 407209 5. clone 407209 5. embryo WhxELS 15. 14.5 Hus musculus CDRA	0517004	9b AA10J385	1.9e-162	86	Mus musculus modiluz. Li Lite fech mouse embryo 13 5dpc 10666014 Mus musculus CDNA clone 554427 5' similar to qb:215030 gnal MyOSIN REGULATORY
0512929	gb[<i>W</i> 75735	3.0e-92	921	Hus musculus meschiz.rl Soares mouse embryo MMMEIl 5 14.5 Hus musculus CUIA					LIGHT CHAIN 2, VENTRICULAR (HUMAN); gb:x65979 M.musculus PLKLC-A mRNA f myosin light chain 2 (MOUSE)
OST2934	дь[н82904	1.8e-75	934	Musiculus Mouse myotonic dystrophy	OST1011	gb AA035805	1.20-98	166	Mus musculus mislal0.rl soares mous embryo NbHEll,5 14.5 Mus musculus c
usr2940	95 4435	1.4e-114	116	Mus musculus mn44cll.rl Dachdinyton mouse embryonic region Mus musculus CDMA clone \$40788 5' similar to gp:x8359 H musculus mRMA for riboromal protein LS. 3'cod	0513017	gb[A2050908	4.8e-123	92%	cone 467226 5' similar to PIN:528237 528237 MADH dehlydrogenate Mus musculus mj21e02.rl Scares mouse embryo (MBRIL) 5 Mt.5 Mus musculus CDM clone 476762 5' similar to SN:APT7 MAT
05T2942	gb w348B2	1.46-91	396	NUS MUSCULUS MC40a05.rl Soures mouse DENNET9.5 Hus musculus CDIA clone	9101730	767180140	2 24-235	200	QOODSO CLATHRIN COAT ASSEMULY PROTE API7 Mus musculus Mouse DNA for small
0512948	gb[AA108292	5.1e-32	811	333300 3 Sattus norvegicus EST0035 rat lambla ZAPII library (C.P.Namel) kattus					GrP-binding protein 510, exon2 and complete cds
				norvegicus cOAA clone pCO9) 5. similar La ADF-ribosylation factor (ARF)-like prorei	21.01.T30	gb U49385	2.16-76		Mus musculus Mus musculus CTP synthetase homolog (CTPSH) mRNA. complete cds
0512953	9090 th qD	1.84-97	984	HUS BUSCULUS ma44dll.rl Soares mouse DJNFF19.5 Nus musculus colls clond DJNFF19.5 nus musculus colls clond	0573035	gb L08651	1.8e-115	1.06	Mus musculus Mus musculus large ribosomal subunit protein mRNA, complete cds
Page Calcula			;	Arpase inhibitor protein precursor, mitochondrial - rat [1]	0ST3037	95606M 116	4.50-34	741	Mus musculus mf84105.rl Soures mouse embryo NDAE11.5 14.5 Mus musculus CDNA

					•				
OST3305 OST3312	9b 088453 9b 078109	1.0c-106 9.7e-59	87.8	Mus musculus Mouse mitta Mus musculus Mus musculus	OST3483	gb X79446	1.40-114	921	Mus musculus M.musculus Odfl mkNA for outer dense fiber protein of sperm
0513323	gb b43643	1.24-132	918	prepro-neurturin mkNA, complete cds Mus musculus Mouse YL-1 mRNA for YL-1	OST3485	95 083824	1.4e-75	198	tails Hope supiens similar to T
0513124	901 X41399	2.26-51	873	protein (nuclear protein with DNA-binding ablility), complete cds Hus austoulus House FSZ mRNA for a	OST3492	81280WJdg	4.7e-139	921	cellispecitic MAL Mus musculus ma08409.rl Scares mouse pJNHYL9.5 Mus musculus cDNA clone
UST 3325	95 028476	6.50-103	941	novel protein Homo sapiens Human mkHA (or KIAA0045	0513494	3313W dy	1.10-138	166	303953 5. Mus musculus mit82d01.rl Soares mouse
osT3349	9b/M18210	2.24-52	948	gune, complete cds Mus musculus Mouse transcription					embryo NuMELLIS 14.5 Mus musculus com clone 174881 5 similar to current nouth plocio curative
0ST3352	gb AA099549	4.9c-63	77.	Tactor S-11. Clone Filt.) Homo sapiens zkôbo4 sl Scares pregnant uterus NbHPU Nomo sapiens					G(1)/G(S)/G(O) GAMAA-5 SUBURIT. (1)
Octra Jak	96) [45] 46	9.1c-69	126	CUNA clone 489679 3' Mus ausculus and 4410 rl Source mouse	00213200	gb[062483	2.1e-180	3 6 4	Mus musculus Kus musculus ubiquitin conjugating enzyme (ubc4) mKNA.
				embryo Numerias 14.5 hus musculus cuma Clone 16871 5: similar to SW:RRCI_MUNAM P25440 RINC3 PROFEIN.[1]	0511501	158650 06	6.8e-54	106	COMPLETE CON 10man feral brain CDHA 5 end GEM-070H03
0513355	91/049185	4.16-40	821	Mus musculus Mus musculus occludin mRNA, complete cds	0513505	gb w40883	J.9e-173	2 66	Mus musculus mc39d07.rl Soares mouse p3NMF19.5 Mus musculus cDNA clone
OST3366	gb AA 122835	2.1e-85	2	Nus musculus marigion. To Beddington mouse embryonic region hus musculus cDNA clone 518900 5°, similar to	0513508	gb H23458	2.0e-119	906	350893 5. Mus.musculus Mus musculus endogenous errovicuslike B-26 (distantly related
				go:Dudgez Cuttin (Horwa); go:Duggz Mouse mRNA for cofflin, complete cds	0513516	gb t14441	5.40-177	\$06	LO Musy/ Lin Rattus norvegicus Rat phosphotidylethanolamine
0213370	gb 567058	4.6e-106	948	Mus sp. Hoxa-4/Hoxa-4 (mice, Genomic, 2556 nt)	OST3517	qb AA015044	5.5e-114	978	N-methyltransferase mRNA, complete cds Mus musculus mh23f10.rl Soares mouse
1711111	9b ₩31107	1.5e-50	711	Homo sapiens 1985ell.rl Soares senescent fibroblasts HUMSF Homo					placenta 4NbMP13.5 14.5 Mus musculus cDNA clone 443371 5'
0Sr3372	gb W64859	2.24-134	166	sapiens CDNA clone 310414 5. Hus musculus me06f10.rl Soares mouse	0573518	95 AA061165	6.3e-99	91 4	Mus musculus mj31f05.rl Soares mouse embryo NDME13.5 14.5 Mus musculus cDNA
				embryo NbwEll.5 14.5 Mus musculus CDNA clone 186711 5 : similar to PIN:A55012 A5011 Signal peptidase 25k chain -	0573521	95/1133756	3.76-70	87	clone 477729 5' similar to TR:E222933 E222933 SUPEROXIDE DISMUTASE Rattus sp. EST110066 Rattus sp. CDNA
0573375	9b AA015237	4.00-44	10.1	dog Mus musculus mh30a10.rl Soares mouse	0573531	gb u19893	6.7e-34	108	 end Rattus norvegicus Rattus norvegicus
-				placents 4MbMP13.5 14.5 Mus musculus	05:1514	ob[032150	5 74-11	9.3	alpha actinin mRNA, complete cds Bos taurus Bos taurus peotide
0513376	gb M27347	4.2e-103	166	Hus musculus Hus musculus p6-5 gene,				;	methionine sulfoxide reductase (msrA)
OS-13388	gb 050264	1.9e-117	186	Nus musculus Mouse mRNA for phospharidy innerity of process of the phospharidy innerity of process of the proce	OST3545	9b н93148	4.0e-103	84%	flox 1.11) dene. complete cds
0513390	gb w34022	3.60-46	1.87	complete cds Mus musculus mb01d09.rl Soares mouse	OST3556	gb W08748	1.9e-129	974	Hus musculus mb48f02.rl Soarce mouse p3NMF19.5 Hus musculus CDHA clone
0511191	011090140	1.76-208	5.6	plamily.5 Mus musculus cDNA clone lleg29 5. Mus musculus Mus musculus Ki anticen	OST355B	95[103386	7.94-132	116	33,200/ 5 Rattus norvegicus Rattus norvegicus (20ne RAHB2-5/8) zinc finger protein
				mkNA, complete cds		and a surface	,	ä	mRNA, J' end cds
0513404	gb A5168895	6.34-109	985	Hus musculus maddg02.rl Life Tech mouse embryo 13 Sdpc 10666014 Hus musculus cDNA clone 614162 5: yimilar	0873561	95/413785	5.1e-64	1 66	Mus musculus may4cil [1 Soares mouse plumF19.5 Mus mausculus cDNA clone 1819356 5. similar to SW:KS27_RAT
ocr3413	7, 81 en jag	3.36-39	116	to go: 12.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.	051.1567	95 44050004	2.8e-48	787	Mus musculus mj39d07.rl Soares mouse embryo NumElls. 5 14.5 Mus musculus CONA
0543425	94 W1116	1.30-105	198	Judos) J Hus mocalus mellids.cl Source mouse embryo NUMELLS 14.5 Mus musculus curn	0511571	95234	2.40-113	914	Mus musculus me'Ja07.rl Boares mouse embryo NME13.5 14.5 Mus musculus CDAA
UST3428	gb[AA189339	3.4e-37	188	Clone Jaybys 5. Mus muculus mc79g04.rl Soures mouse lymph node NUHLN Hus musculus CDNA	osr1575	yb[AA080212	6.00-90	116	CLORE SYLING 3' Hus musculus mj99006.rl Soares mouse plant19.5 Mus musculus cDNA clone 484210 5' wimilar to ob:x88079 5-100
OSF3441	958155 06	7.90-66	77.1	Mus sp. Mos gene fmice, embryos,	0.54150	cts \$74622	110-19	76%	PROTEIN, ALPHA CHAIN (HUNAN)
0513450	gb x58426	7.1e-53	196	Huses, the second of the secon				}	phosphatase IM 21 kda regulatory submit Chickens dizzard smooth
0273457	9D[W87064	9.04-166	11.6	embryo Ribelli. S. 14.5 mus musculus ciril	0:13582	gb x74.150	1.5e-74	164	muscle, minA, 1598 nt. Hus musculus H.musculus XPAC Kurderma
OST3460	ob[AA185213	4.7e-134	106	clone 418521 5'		at the desired	47,1	5	Pigmentosum group A Correcting gene, exon 6
				Lymph note Nights for musically curv clove 642974 5: similar to TR: E241948 F241948 CHRANGOME UTI HEADIN: FHAME	Participan	u concellat.			nomo saprena noman prantina nacisoria: sechange factor pbJ2 mRMA, complete cds
0573480	ab AA 118567	9.46-100	3.63	ORF VCLOS4C. Nus musculus mnl2a08.rl Beddington	UST3602	95 K15062	2.3e-107	306	Homo sapiens yf86h05.rl Homo sapiens cDNA clone 29481 5.
				mouse embryonic region Mus musculus	0ST3604	gb M22756	4.94-119	844	Rattus norvegicus Rat 24-kDa subunit of mitochondrial NADM dehydrogenase
OST3481	30 x56906	1.00-121	951	Mus musculus Mouse OP-1 mRHA for ostwogenic protein 1	05T360H	96/034994	5.44-101	158	MRNA, 3' end Homo sapiens fluman 046-dependent

9091230	ob14A165901	2.40-129	196	procedn kinase catalytic subunit (DRA-PECS) may, complete cds Hur susculla mr73c01 rl Scare: souse	0513788	88 gb AA014426	9.70-55	101	Mus musculus mg64b01.rl Soares mouse embryo NDHE13.5 14.5 Mus musculus CDNA clone 43967 5. similar to
				lymph node NbMLR Mus musculus class	_				SWINDTH BOVIN 002367 NADH-UBIQUINGNE OXIDOREDUCTASE 017 SUBURIT
UST36.11	gb AA028590	2.1e-152	97.	Mus musculus mi21c12.rl Soares mouse	OST3789	89 gb/b13544	9.50-67	37.6	Mus musculus Mouse mRNA for primase
				Clone 464182 5' similar to WP:R01M2.6	OST3807	07 gb[w26968	3.8e-51	804	Homo sapiens 1617 Human retina CONA randomly primed sublibrary Homo
0273642	gb[k85211	1.46-47	74%	Homo sapiens yodidil.si Homo sapiens CDM cione 180501 3: similar to	STALTSO	18 (D) M28248	3.Be-48	963	sapiens cDNA unidentified cloning vector Moloncy
				SP.S19586 S19586 N-HETHYL-D-ASPARTATE BECERMON CLITTAMATE-BINDING CHAIN	-				murine leukemia virus retroviral
05TJ645	gb H14951	1.3e-104	106	Must musculus Mouse insulin-like growth	OST3819	19 95 155632	3.80-35	91.	Homo sapiens yb39b03.rl Homo sapiens
0573647	95 014721	1.7e-36	761	MAGGOI II (ICF-11) BRNA, COMPTETE COS MAG MAGGOLIAS AUGGOTIAS C-101	.ca(+->c	Oraganation to	1 7	4	SP: 7205.10 CE00629
	224160441.45	1 4.1.109		partial cds			10.57		Heart Mailli DW Home Supiens CDNA Clone
	a received in fi		;	placenta MbMP1.5 14 5 Hus muculus CDNA clone 455981 5' similar to	0513831	77.07W dg 18	J. Se-121	166	Mus musculus med4a02.rl Soares mouse embryo NbME13.5 14.5 Mus musculus CDN
osr3652	gb s60494	3.14-31	941	SW.AAP_HUMKN QO4941 INTESTINAL MEMBKANE AR PROTEIN. [1] Hus sp. gamma-phosphorylase kinare	0513839	39 gb H86008	1.4e-103	861	Homo sapiens EST02533 Homo sapiens CONA clone HPUCK19 similar to
				(alternatively spliced) (mice, muscle, balb/C, Genomic, 4204 nt. segment 4 of 41	0513843	43 gb 282190	2.8e-51	88	Hypotherical 43.5K procein Homo sepiens Human DNA sequence EEQUENCING IN PROGRESS from clone
OST 1662	gb(037427	3.16-204	196	Rattus norvegicus Rattus norvegicus phospholipid hydroperoxide glutathione	0013849	49 gb W64986	1.3e-173	876	180M12; HTCS phase 1 Mus musculus mc04c05.rl Source mouse
0311669	gb W55918	3.0e-35	864	peroxidase mRNA, complete cus Homo sapiens zc01f12.s1 Soares parathyroid tumor NbHPA Homo sapiens					embry Nowells at 4.5 Mbs musculus cuma clone 186504 5: similar to SW:VSH7_DICDL P14327 VEGETATIVE
OST3681	96 455833	7.64-94	938	CDNA clone 121261 3' similar to WP:EO4f6_2 CE01214 Hus musculus md07b01.rl Soares mouse	OST3851	51 954US1037	1.04-135	841	SPECIFIC PROTEIN H7. [1] Mus musculus Hus musculus [1]-inc-finger transcription factor
				embryo NbHEll.5 14.5 Mus musculus cDNA clone 167657 5' similar to gb:U37874	0ST3858	58 gb[x56135	4.76-237	1/6	(CTCF) mRNA, complete cds Hus musculus Mouse mkNA for
0573694	gb W38194	5.44-71	934	House rekn gene: (Mouse) Hose sapiens zelse05:s Source	05T38C4	64 gb b19493	9.8e-33	156	Mus musculus Mouse 3directed CDMA, wiscendrike
0211100	ub AA038243	4.94-171	166	CDNA clone 122400 3. Hus musculus mi82408.rl Scares mouse	0573869	69 gb W41525	4.40-100	85%	Hus musculus mc45b04.rl Soares mouse planf19.5 Mus musculus cunA clone
	<u>.</u>			plinkF19.5 Mus musculus CORA clone 473103 5: similar to SM:SARL_RABIT	0513897	97 gb[w10485	3.80-97	356	351439 5. Mus musculus ma53e06.rl Source mouse
0513703	gb[w47847	7.Be-71	82.8	FY231 SANCOLFIA: 111 Mus musculus me82f12.11 Soures mouse embryo NDME13.5 14.5 Hos musculus CDNA	0513903	03 gb[w59388	1.26-108	861	114434 5' Mus musculus md79f02.rl Soares mouse
OCT1704	gb AA048648	4.6e-68	166	clone 155055 5. Hus musculus mjlla07.rl Soares mouse embryo NDME11.5 14.5 Mus musculus CURA					clone 374619 5' similar to gb:U07151 ADP-RIBOSYLATION FACTOR-LIKE FROTEIN 3
UST3708	gb[AA002275	7.44-89	97.6	clone 47876 5. Hus masculus my41h01.fl Source mouse embryo (HDHE115 14 5 Hus musculus CHIA embryo (ACETA)	05T1905 05T1909	05 95 05 05 000 000 000 000 000 000 000	8.0c-102 1,2c-80	921	HUBBAN) Mus musculus Mouse Murri mKWA, exen Mus musculus mi61a06.rl Seares mouse Alacata Amanal 6 14 6 mus musculus
911f1350	gb AAU34685	H. 2e-119	106	PHOSPINATION SITE OF SUPPLY OF WARD WAS MAKED WAS MASSED OF SOURCES MOUSE OF SUPPLY OF	OST1917	17 90 244044	B.7e-81	871	CONA clone 455410 5' Homo sapiens H. sapiens partial CUNA
				embryo NDME13.5 14.5 Mus musculus cDNA clone 467587 5' similar to gb:L19527 605 Kludschal, PROTEIN L27	0513924	24 gb J04699	3.9e-32	841	sequence; clone c-irco/ Mus musculus Mouse nicotinic acetylcholine receptor beta subunit
0573729	419303	2.90.97	85.8	Homo sapiens 1025402.rl Source fetal lung Whit1DW Homo sapiens cuth clone 101075 5	0573925	25 gb[W23511	1.20-88	191	INACINE) genu, complete eds Homo sapiens zb46e02.rl Soarus fetul Lung NHLLISW Homo sapiens cDNA clone
11,71,130	3051M1305	1.3e-131	376	HUS MUSCULUS MANUTOD. IL SOJEES MOUSU p)WHF19.5 HUS MUSCULUS CUMA Clone]17051 5' similar to SW:PRCF.HUWAN Pq)306 PROTENSOME COMPONERT MELL-1	0013931	st gb[U14957	1.64-36	118	Juneau 5. Homo supiens Human 51K isoform of Type II phosphatidylinusitol-4-phosphate 5-kinase (PIPK) mRNA, complete cds.
est 7 LTso	95 (45014575	5,20-100 971	17.4	HKEUN:OR Hus musculou micflgol, Il Soares mouse enbryo (1981), 5, 14, 5 Hus musculus CDIA	OST 3945	45 gistation4	1 . 6c- 122	11.6	Mus musculus mb25c09.rl Scares mouse plump19.5 Hus musculus conn clone 310448 5'
7.27.ET.20	95/W77924	2.64-99	¥CB	Clone (66815 5 : stails TO C EN SYNY, VEAT P. 1911 A ACTIVIL-THIN SYNTHETISE, MITOCHONUMIAL PRECURSON Home spainen 2471104, II Soares fectal heart NDHHIJSW Homo sapiens CDNA clone	rangaga	57 gb AA051293	2.8v-14)	196	ukus musculus m400hl0.r.1 Soares mouse embryo NbHEll 5 14 5 Hus musculus cDNA clone 478627 5. similar to swifts.humu P2064 FRANSFORHING GROWTH FACTOR BETA-1 BINDING PROTEIN
0ST3759	9b x64840	7.66-51	97.4	346111 5. Hus musculus M.musculus ALP1 mRNA Huma esate Human chicants citat	0513960	50 gb D38614	1.1c-88	821	PRECURSOR HUS BUSCULUS MOUSE 921-5 BRNA for PRESENDANTIC CROSSING COMMISSION COM
0513775	95/018282	1.64-57	97.	form musculus House 1'-directed cuth.	1961150	51 95 067988	6.6e-37	111	Homo sapiens Human guanylate kinace associated protein (GKAP) mRUA.

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Mus marculus mid7ell.rl Soares mouse embryo NMRELLS.54. Hws marculus CDMA clone 479276 5. similar to qb:UJ1705 Juna marculus demeticus CSTBL/63 places a gluethione (HOUSE) in Soares souse Diagna gluethione (HOUSE) al Homo sapient (HOUSE) rl Soares mouse lux marculus mid7blo.lr 15 oares mouse placenta 4MBMP115 14.5 Homo sapient CDMA clone 455815 4.5 Huss marculus SW:A4 HUAM Q04941 INTESTIMA. HORSENDE 4 PROTECTIVA. (1) flucobate (1977212) Homo sapients cDMA clone 529412 3. similar to cDMA clone 529412 14.5 Hus marculus cDMA clone 529412 3. 44.5 Huam anscrulus millado4.rl Soares mouse cabryo NEMEL) 3.14.5 Hus marculus CDMA clone 187950 5. 4.4 Huam anscrulus millado1.rl Soares mouse cDMA clone 442213 5. similar to cDMA clone 442213 5. similar to human Hus marculus millado1.rl Soares mouse CDMA clone 442213 5. similar to human Hus marculus millado1.rl Soares mouse cDMA clone 442213 5. similar to human Hus marculus millado1.gene, partial cds and sucrulus deman (TDMA) gene, partial cds and sucrulus (TDMA) gene, complete

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Figure 8 cont'd.

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	2.70-89	9.3e-205	4.8e-70	3.06-173			2.36-40			00.00	20.0	1.5e-115			8.7e-154		2.4e-82		4. Je-169		4.06-40		8.96-52		1.4e-135			6.44-62	2.0e-127																		
•	gb AA203787	90 551016	qb z31263	gb W53187	2		gb (AA048921			216011142	9170111 06	gb AA023146			gb AA070774		gb[WS4737		gb AA013789		96 016175		gb1AA007519		gb AA000024			gb H18210	gb J04696																		
	0514223	OST4228	OST4229	OST4235			OST4243			346,400		OST4247			OST4251		OST4254	-	0514258		OST4281		OST4283		OST428B			OST4315	OST4319																		
embryo NDME13.5 14.5 Mus musculus CDNA	Nus musculus Mouse serum amyyloid A bseudodene (psi-SAA)	Homo sapiens yf31a08.sl Homo supiens CDMA clone 128630 3:	Mus musculus mg34e07.rl Soares mouse	close 425700 5:	myeloid secondary granule protein mRNA	Mus musculus Mus musculus expressed	Homo sapiens Human mRNA for KIAA0280	gene, partial cds	neuron (#937233) Nono sapiens CD4A	clone 546559 J' similar to TR:G600529	SUBUNIT	Homo sapiens H. sapiens partial cDNA	sequence; clone c-lad08	plinkije, S Mus musculus cDNA clone	Mus musculus mb76g12.rl Soares mouse	plNMF19.5 Mus musculus cDNA clone	Rattus norvegicus R.norvegicus mRNA	Rattus norvegicus Rat mRNA for	dihydropyrimidinase, complete cds	eabryo NDMEI3.5 14.5 Mus musculus CDMA	Clone 400594 5' Mus musculus mb96g01.rl Soares mouse	plinfile.5 Mus musculus cDNA clone	Homo sapiens zk54h03.s1 Soares	CDNA clone 486677 3'	Rattus sp. EST103504 Mattus sp. cumA	Hus musculus me27f01.rl Soares mouse embryo NbME13.5 14.5 Mus musculus cDNA	clone 388729 5' similar to	KD PROTEIN IN PETIL2-ILSI INTERCENIC	Rattus norvegicus similar to none	Hus musculus Mouse many for prothymosin alpha	Mus musculus md08h09.rl Soares mouse	clone 167841 S' similar to PIR:A56059	Asousy procein-cyrosine-phosphatase Mus musculus Mus musculus	transcription factor TFEB mRNA,	partial cus Mus musculus Mouse mRNA (clone	lambda-16) for hypothetical protein A Rattus norvegicus Rat mRNA for lens	betaB1-crystallin (pRLbeta B1-3)	Rattus norvegicus Rattus norvegicus	partial cds	Mus musculus Mus musculus X inactive specific transcribt (Xist) pene.	cosmid MB4-14A, fragment 2	Mus musculus mf49h12.rl Soares mouse	eabryo NDME13.5 14.5 Mus musculus cDRA	CIONE 408455 5' SIMILAR TO SW:GLYM_HUMAN P34897 SERINE	HYDROXYMETHYLTRANSFERASE,	MITOCHONDRIAL Mus musculus mc3le07.rl Soares mouse	DJWHI19.5 Mus musculus cDNA clone
:	106	821	196	:	1.	941	921		000			861	474	;	106		916	198			961		169	į	1 C R	916			894	Š	911		196		196	851		906		8		821				871	;
:	2.6e-111	4.7e-45	1.9e-112	101-101	171-96-7	2.0e-155	7.5e-93		1.28-34			7.6e-63	110-118	91.5	6.0e-115		2.0e-105	3.3e-140			6.54-90		2.4e-33	;	1.04-84	3.7e-121			5.7e-74	4.46-41	1.5e-133		2.6e-111		1.3e-161	1.50-58		8.0e-169		1.3e-38	1	2.2e-83				8.94-38	
	gb H13524	gb R16778	gb AA000314	-41.12262	/67/17/q6	gb L26664	ab D87470	1000000	gb AA084 / 04			gb F03500	-1-1430610	oreoculos.	gb W36515		gb x82021	gb D63704	100000	Book / Milab	gb w20730		gb AA044274		gb[H31489	gb w71052			gb C07091	gb x56135	gb W54510		ob1036393		ab x56046	DOS SON JAPO	100000	gb U53859	,	gb U41395	;	ob w85357				ob W34635	200100
	0513988	OST1993	OST4002		0214003	0574011	OST4028		0574033			OST4051	. ,	1905.150	0574070		0514073	OST4074		0214108	0574114		osr4131		0574134	OST4140			OST4142		OST4148		OST4149		OST4154	00000166		OST4166		OST4174		OST4191				OST4194	

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IPC(6) :0	SIFICATION OF SUBJECT MATTER C12Q 1/68; C12N 5/02, 5/06, 15/00, 15/64; C07H 35/6, 320.1, 325, 357; 536/23.1, 24.2; 800/2		
According to	International Patent Classification (IPC) or to both	national classification and IPC	
	OS SEARCHED		
	cumentation searched (classification system followed	by classification symbols)	
	35/6, 320.1, 325, 357; 536/23.1, 24.2; 800/2		
Documentati	on searched other than minimum documentation to the	extent that such documents are included	in the fields searched
Electronic d	ata base consulted during the international search (na DIALOG	me of data base and, where practicable	, search terms used)
C. DOC	UMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.
X	SAUER, B. Site-specific recomb applications. Current Opinion in Biotect pages 521-527, see the entire article.	ination; developments and chnology. May 1994, Vol. 5,	1-8, 10, 20 and 28
Y	SEKINE et al. Frameshifting is required transposase encoded by insertion sequentus. June 1989, Vol. 86, pages "Frameshifting in Other Systems", pages	nce 1. Proc. Natl. Acad. Sci. 4609-4613, see especially	10
х	WANG, et al. High frequency recomb human chromosomes mediated by an Cre recombinase. Somatic Cell and M 1996, Vol. 21, No. 6, pages 429-441,	adenovirus vector expressing olecular Genetics. 09 March	8
X Furth	ner documents are listed in the continuation of Box C	See patent family annex.	
<u> </u>	ecial categories of cited documents:	"T" later document published after the int	ernational filing date or priority
·A· do	cument defining the general state of the art which is not considered	date and not in conflict with the app the principle or theory underlying the	invention
1	be of particular relevance rlier document published on or after the international filing date	"X" document of particular relevance; the considered novel or cannot be considered.	e claimed invention cannot be
1. do	cument which may throw doubts on priority claim(s) or which is ed to establish the publication date of another citation or other	when the document is taken alone	
sp.	ed to establish the publication date of another cleanon of other secial reason (as specified) cument referring to an oral disclosure, use, exhibition or other	"Y" document of particular relevance; the considered to involve an inventive combined with one or more other suc-	step when the document is
me	cument reterring to an oral disclosure, use, exhibition of other seans coment published prior to the international filling date but later than	being obvious to a person skilled in document member of the same paten	the art
the	actual completion of the international search	Date of mailing of the international se	
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Commissio Box PCT	mailing address of the ISA/US ner of Patents and Trademarks n, D.C. 20231 lo. (703) 305-3230	Authorized officer WILLIAM SANDALS Telephone No. (703) 308-0196	abor

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C (Continue	ation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevan	t passages	Relevant to claim No
Y	ODELL et al. Site-directed recombination in the genome transgenic tobacco. Molecular and General Genetics. 11 1990, Vol. 223, pages 369-378, see especially Figure 1 a "Result" section.	October	1-8, 10, 20
x	DYMECKI, S. A modular set of Flp, FRT and LacZ fus vectors for manipulating genes by site-specific recombinations of Gene. 01 June 1996, Vol. 171, pages 197-201, see esperigure 1.	ation.	10
X	HAAS et al. TnMax - a versatile mini-transposon for the of cloned genes and shuttle mutagenesis. Gene. 11 Aug Vol. 130, pages 23-31, see especially the abstract.		8
Y	WO 88/01646 (ALLELIX INC.) 10 March 1988 (10.10. especially pages 1-3.	88), see	1-8, 10 and 20

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Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
Please See Extra Sheet.
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. X No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-8, 10, 20 and 28
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

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BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1.

Group I, claim(s) 1-7, 8, 10, 20 and 28, drawn to a library of cultured eucaryotic cells made by a process comprising treating a group of cells with a vector that mediates the splicing of a foreign exon internal to a cellular transcript, the use of the cell from the library to generate a non-human transgenic animal, and the method of making the cell comprising the vector and the use of the vector to make the library of cultured eukaryotic cells.

Group II, claim(s) 9, 11-18, drawn to a vector construct for replacing the 3' end of an animal cell transcript with a foreign exon.

Group III, claim(s) 19, 21 and 22, drawn to the use of a vector according to claim 9. Group IV, claim 23, drawn to a stably transduced animal cell that incorporates the vector of claim 16.

Group V, claims 24-27, drawn to a method of altering a region of DNA by adding or deleting DNA.

The inventions listed as Groups I-V do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following teasons: the first group contains the product, a library of cultured eukaryotic cell, a method of using the cells to produce a non-human transgenic animal and a method of making the cells. The additional groups are directed to different vectors having different compositions than the vector used in the first group, cell lines containing those vector constructs and methods of altering the cellular genome. The first group contains a vector having a different composition than the other vectors and therefore the special technical feature present in the first group does not occur in the other groups.